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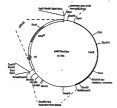
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(54) Title: IN VIVO HOMOLOGOUS SEQUENCE TARGETING IN EUKARYOTIC CELLS



(57) Abstract

The invention relates to methods for targeting an exogenous polynacleotide or exogenous complementary polynacleotide pair to a predetermined endogenous DNA target sequence in a cokaryotic cell by homologous pairing, particularly for aftering an endogenous DNA requence, such as a chromosomal DNA sequence, typically by targeted homologous recombination. In certain embodiments, the invention relates to methods for targeting an exogenous polymaclocoide having a linked chemical substituent to a produtermined endogenous DNA sequence in a metabolically active enkaryotic cell, generating a DNA sequence-specific targeting of one or more chemical substituents in an intact nucleus of a metabolically active sukeryotic cell, generally for purposes of aftering a predetermined endogenous DNA sequence in the cell. The invention also relates to compositions that contain exogenous targeting polynucleotides, complementary pairs of exogenous targeting polynucleotides, chemical substituents of such polynucleotides, and recombinase proteins used in the methods of the invention.

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IN VIVO HOMOLOGOUS SEQUENCE TARGETING IN EUKARYOTIC CELLS

FIELD OF THE INVENTION

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The invention relates to methods for targeting an exogenous polynucleotide or exogenous complementary polynucleotide pair to a predetermined endogenous DNA target 10 sequence in a eukaryotic cell by homologous pairing, particularly for altering an endogenous DNA sequence, such as a chromosomal DNA sequence, typically by targeted homologous recombination. In certain embodiments, the invention relates to methods for targeting an exogenous polynucleotide having a linked chemical substituent to a predetermined endogenous DNA sequence in a metabolically active eukaryotic cell, generating a DNA sequence-specific targeting of one or more chemical substituents in an intact nucleus of a metabolically active eukaryotic cell, generally for purposes of altering a 20 predetermined endogenous DNA sequence in the cell. The invention also relates to compositions that contain exogenous targeting polynucleotides, complementary pairs of exogenous targeting polynucleotides, chemical substituents of such polynucleotides, and recombinase proteins used in the methods 25 of the invention.

BACKGROUND

Homologous recombination (or general recombination) is defined as the exchange of homologous segments anywhere along a length of two DNA molecules. An essential feature of general recombination is that the enzymes responsible for the recombination event can presumably use any pair of homologous sequences as substrates, although some types of sequence may be favored over others. Both genetic and cytological studies have indicated that such a crossing-over process occurs between pairs of homologous chromosomes during meiosis in higher organisms.

Alternatively, in site-specific recombination, exchange occurs at a specific site, as in the integration of

40 phage λ into the E. coli chromosome and the excision of λ DNA

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from it. Site-specific recombination involves specific sequences of the phage DNA and bacterial DNA. Within these sequences there is only a short stretch of homology necessary for the recombination event, but not sufficient for it. The enzymes involved in this event generally cannot recombine other pairs of homologous (or nonhomologous) sequences, but act specifically on the particular phage and bacterial sequences.

Although both site-specific recombination and homologous recombination are useful mechanisms for genetic engineering of DNA sequences, targeted homologous recombination 10 provides a basis for targeting and altering essentially any desired sequence in a duplex DNA molecule, such as targeting a DNA sequence in a chromosome for replacement by another sequence. Site-specific recombination has been proposed as one 1.5 method to integrate transfected DNA at chromosomal locations having specific recognition sites (O'Gorman et al. (1991) Science 251: 1351; Onouchi et al. (1991) Nucleic Acids Res. 19: 6373). Unfortunately, since this approach requires the presence of specific target sequences and recombinases, its utility for targeting recombination events at any particular chromosomal location is severely limited in comparison to targeted general recombination.

For these reasons and others, targeted homologous recombination has been proposed for treating human genetic 25 diseases. Human genetic diseases include: (1) classical human genetic diseases wherein a disease allele having a mutant genetic lesion is inherited from a parent (e.g., adenosine deaminase deficiency, sickle cell anemia, thalassemias), (2) complex genetic diseases like cancer, where the pathological 30 state generally results from one or more specific inherited or acquired mutations, and (3) acquired genetic disease, such as an integrated provirus (e.g., hepatitis B virus). However, current methods of targeted homologous recombination are inefficient and produce desired homologous recombinants only 35 rarely, necessitating complex cell selection schemes to identify and isolate correctly targeted recombinants. A primary step in homologous recombination is DNA .

strand exchange, which involves a pairing of a DNA duplex with

at least one DNA strand containing a complementary sequence to form an intermediate recombination structure containing heteroduplex DNA (see, Radding, C.M. (1982) Ann. Rev. Genet. [4: 405; U.S. Patent 4,888,274). The heteroduplex DNA may take several forms, including a triplex form wherein a single complementary strand invades the DNA duplex (Heich et al. (1990) Genes and Bevelogment &: 1951) and, when two complementary DNA strands pair with a DNA duplex, a classical Holliday recombination joint or chi structure (Holliday, R. 10 (1944) Genet. Rem. 5: 282) may form, or a double-D loop

("Diagnostic Applications of Double-D Loop Pormation" U.S.S.N 07/755,462, filed 4 September 1991, which is incorporated herein by reference). Once formed, a heteroduplex structure may be resolved by strand breakage and exchange, so that all or a portion of an invading DNA strand is spliced into a recipient DNA duplex, adding or replacing a segment of the recipient DNA duplex. Alternatively, a heteroduplex structure may result in gene conversion, wherein a seguence of an invading strand is transferred to a recipient DNA duplex by repair of mismatched bases using the invading strand as a template (Egamss. 3rd Ed.

20 bases using the invading strand as a template (Ganss, 3rd Ed. (1987) Levin, B., John Wiley, New York, NY; Lopes et al. (1987) Nucleic Acide Rass. Lis. 5643). Whether by the mechanism of breakage and rejoining or by the mechanism(s) of gene conversion, formation of heteroduplex DNA at homologously paired joints can serve to transfer genetic sequence

information from one DNA molecule to another.

The ability of homologous recombination (gene conversion and classical strand breakage/rejoining) to transfer genetic sequence information between DNA molecules makes

genetic sequence information between DNA molecules makes: 30 targeted homologous recombination a powerful method in genetic engineering and gene manipulation.

The ability of mammalian and human cells to incorporate exogenous genetic material into genes residing on chromosomes has demonstrated that these cells have the general senzymatic machinery for carrying out homologous recombination required between resident and introduced sequences. These targeted recombination events can be used to correct mutations at known sites, replace genes or gene segments with defective

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ones, or introduce foreign genes into cells. The efficiency of such gene targeting techniques is related to several parameters: the efficiency of DNA delivery into cells, the type of DNA packaging (if any) and the size and conformation of the 5 incoming DNA, the length and position of regions homologous to the target site (all these parameters also likely affect the ability of the incoming homologous DNA sequences to survive intracellular nuclease attack), the efficiency of recombination at particular chromosomal sites and whether recombinant events 10 are homologous or nonhomologous. Over the past 10 years or so, several methods have been developed to introduce DNA into mammalian cells: direct needle microinjection, transfection, electroporation, retroviruses, adenoviruses, and other viral packaging and delivery systems, liposomes, and most recently 15 techniques using DNA-coated microprojectiles delivered with a gene gun (called a biolistics device), or narrow-beam lasers (laser-poration). The processes associated with some types of gene transfer have been shown to be both mutagenic and carcinogenic (Bardwell, (1989) Nutagenesis 4:245), and these possibilities must be considered in choosing a transfection approach.

The choice of a particular DNA transfection procedure depends upon its availability to the researcher, the technique's efficiency with the particular chosen target cell 25 type, and the researchers concerns about the potential for generating unwanted genome mutations. For example, retroviral integration requires dividing cells, most often results in nonhomologous recombination events, and retroviral insertion within a coding sequence of nonhomologous (i.e., non-targeted) gene could cause call mutation by inactivating the gene's 30 coding sequence (Friedmann, (1989) Science 244:1275). Newer retroviral-based DNA delivery systems are being developed using defective retroviruses. However, these disabled viruses must be packaged using helper systems, are often obtained at low 35 titer, and recombination is still not site-specific, thus recombination between endogenous cellular retrovirus sequences and disabled virus sequences could still produce wild-type retrovirus capable of causing gene mutation. Adeno- or polyona

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virus based delivery systems appear very promising (Samulski et al., (1991) EMBO J. 10: 3941: Gareis et al., (1991) Cell. Molec. Biol. 37: 191: Rosenfeld et al. (1992) Cell 68: 143) although they still require specific cell membrane recognition 5 and binding characteristics for target cell entry. Liposomes often show a narrow spectrum of cell specificities, and when DNA is coated externally on to them, the DNA is often sensitive to cellular nucleases. Newer polycationic lipospermines compounds exhibit broad cell ranges (Behr et al., (1989) Proc. 10 Natl. Acad. Sci. USA 86:6982) and DNA is coated by these compounds. In addition, a combination of neutral and cationic lipid has been shown to be highly efficient at transfection of animal cells and showed a broad spectrum of effectiveness in a variety of cell lines (Rose et al., (1991) BioTechniques 15 10:520). Galactosylated bis-acridine has also been described as a carrier for delivery of polynucleotides to liver cells (Haensler JL and Szoka FC (1992), Abstract V211 in J. Cell. Biochem. Supplement 16F, April 3-16, 1992, incorporated herein by reference). Electroporation also appears to be applicable 20 to most cell types. The efficiency of this procedure for a specific gene is variable and can range from about one event per 3 x 104 transfected cells (Thomas and Capecchi, (1987) Cell 51:503) to between one in 107 and 108 cells receiving the exogenous DNA (Koller and Smithles, (1989) Proc. Natl. Acad. 25 Sci. (U.S.A.) 86: 8932). Microinjection of exogenous DNA into the nucleus has been reported to result in a high frequency of stable transfected cells. Zimmer and Gruss (Zimmer and Gruss (1989) Nature 338: 150) have reported that for the mouse hox1.1 gene, 1 per 150 microinjected cells showed a stable homologous site specific alteration.

Several methods have been developed to detect and/or select for targeted specific recombinants between vector DNA and the target homologous chromocomal sequence (ggg, Capecchi, (1989) Science 244:1288 for review). Cells which exhibit a specific phenotype after specific homologous recombination, such as occurs with alteration of the hert gene, can be obtained by direct selection on the appropriate growth medium. Alternatively, a selective marker sequence such as nge can be

incorporated into a vector under promoter control, and successful transfection can be scored by selecting G418^r cells followed by PCR to determine whether neo is at the targeted site (Joyner et al., (1989) Nature 338:153). A positivenegative selection (PNS) procedure using both neo and HSV-tk genes allows selection for transfectants and against nonhomologous recombination events, and significantly enriched for desired disruption events at several different mouse genes (Mansour et al., (1988) Nature 336:348). This procedure has the advantage that the method does not require that the 10 targeted gene be transcribed. If the targeted gene is transcribed, a promoter-less marker gene can be incorporated into the targeting construct so that the gene becomes activated after homologous recombination with the target site (Jasin and 15 Berg, (1988) Genes and Development 2:1353; Doetschman et al. (1988) Proc. Natl. Acad. Sci. (U.S.A.) 85: 8583; Dorini et al., (1989) Science 243:1357; Itzhaki and Porter, (1991) Nucl. Acids Res. 19:3835). Recombinant products produced using vectors . with selectable markers often continue to retain these markers 20 as foreign genetic material at the site of transfection, although loss does occur. Valancius and Smithies (Valancius and Smithies, (1991) Molec. Cellular Biol. 11:1402) have recently described an "in-out" targeting procedure that allowed a subtle 4-bp insertion modification of a mouse hort target 25 gene. The resulting transfectant contained only the desired modified gene sequence and no selectable marker remained after the "out" recombination step. Cotransformation of cells with two different vectors, one vector containing a selectable gene and the other used for gene disruption, increases the 30 efficiency of isolating a specific targeting reaction (Reid et al., (1991) Molec. Cellular Biol. 11:2769) among selected cells that are subsequently scored for stable recombinants. Unfortunately, exogenous sequences transferred into

eukaryotic cells undergo homologous recombination with
homologous endogenous sequences only at very low frequencies,
and are so inefficiently recombined that large numbers of cells
must be transfected, selected, and screened in order to
generate a desired correctly targeted homologous recombinant

(Kucherlapati et al. (1984) Proc. Natl. Acad. Sci. (U.S.A.) 81: 3153; Smithies, O.(1985) Nature 317; 230; Song et al. (1987) Proc. Natl. Acad. Sci. (U.S.A.) 84: 6820: Doetschman et al. (1987) Nature 330: 576: Kim and Smithies (1988) Nucleic Acids 5 Res. 16: 8887; Doetschman et al. (1988) op.cit.; Koller and Smithies (1989) op.cit.: Shesely et al. (1991) Proc. Natl. Acad. Sci. (U.S.A.) 88: 4294; Kim et al. (1991) Gene 103: 227. which are incorporated herein by reference).

10 Recently, Koller et al. (1991) Proc. Natl. Acad. Sci. (U.S.A.) 88: 10730 and Snouwaert et al. (1992) Science 257: 1083, have described targeting of the mouse cystic fibrosis transmembrane regulator (CFTR) gene for the purpose of inactivating, rather than correcting, a murine CFTR allele. 15 Koller et al. employed a large (7.8kb) homology region in the targeting construct, but nonetheless reported a low frequency for correct targeting (only 1 of 2500 G418-resistant cells were correctly targeted). Thus, even targeting constructs having . long homology regions are inefficiently targeted.

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Several proteins or purified extracts having the property of promoting homologous recombination (i.e., recombinase activity) have been identified in prokaryotes and eukaryotes (Cox and Lehman (1987) Ann. Rev. Biochem. 56:229; Radding, C.M. (1982) op.cit.; Madiraju et al. (1988) Proc. 25 Natl. Acad. Sci. (U.S.A.) 85: 6592; ; McCarthy et al. (1988) Proc. Natl. Acad. Sci. (U.S.A.) 85: 5854; Lopez et al. (1987) op.cit., which are incorporated herein by reference). These general recombinases presumably promote one or more steps in the formation of homologously-paired intermediates, strand-30 exchange, gene conversion, and/or other steps in the process of homologous recombination.

The frequency of homologous recombination in prokaryotes is significantly enhanced by the presence of recombinase activities. Several purified proteins catalzye 35 homologous pairing and/or strand exchange in vitro, including: E. coli recA protein, the T4 uvsX protein, and the rec1 protein from Ustilago maydis. Recombinases, like the recA protein of E. coli are proteins which promote strand pairing and exchange.

The most studied recombinase to date has been the recA recombinase of E. coli, which is involved in homology search and strand exchange reactions (see, Cox and Lehman (1987) op.cit.). RecA is required for induction of the SOS repair 5 response, DNA repair, and efficient genetic recombination in E. coli. RecA can catalyze homologous pairing of a linear duplex DNA and a homologous single strand DNA in vitro. In contrast to site-specific recombinases, proteins like recA which are involved in general recombination recognize and promote pairing of DNA structures on the basis of shared homology, as has been shown by several in vitro experiments (Hsieh and Camerini-Otero (1989) J. Biol. Chem. 264: 5089: Howard-Flanders et al. (1984) Nature 309: 215; Stasiak et al. (1984) Cold Spring Harbor Symp. Quant. Biol. 49: 561; Register et al. (1987) J. Biol. Chem. 15 262: 12812). Several investigators have used reck protein in vitro to promote homologously paired triplex or other threestranded joint DNA molecules (Cheng et al. (1988) J. Biol. Chem. 263: 15110; Ferrin and Camerini-Otero (1991) Science 354: 1494: Ramdas et al. (1989) J. Biol. Chem. 264: 17395; Strobel 20 et al. (1991) Science 254: 1639; Hsieh et al. (1990) op.cit.; Rigas et al. (1986) Proc. Natl. Acad. Sci. (U.S.A.) 83: 9591; and Camerini-Otero et al. U.S. 7,611,268 (available from Derwent), which are incorporated herein by reference). Unfortunately, many important genetic engineering manipulations 25 involving homologous recombination, such as using homologous recombination to alter endogenous DNA sequences in a living cell, cannot be done in vitro. Further, gene therapy requires highly efficient homologous recombination of targeting vectors with predetermined endogenous target sequences, since 30 selectable marker selection schemes such as those currently available in the art are not usually practicable in human

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beings. Thus, there exists a need in the art for methods of efficiently altering predetermined endogenous genetic sequences by homologous pairing and homologous recombination in vivo by introducing one or more exogenous targeting polynucleotide(s) that efficiently and specifically homologously pair with a predetermined endogenous DNA sequence. There exists a need in

the art for high-efficiency gene targeting, so that complex in vitro selection protocols (e.g., neo gene selection with G418) which are of limited utility for in vivo gene therapy on affected individuals, are avoided.

SUMMARY OF THE INVENTION

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It is an object of the present invention to provide methods for targeting an exogenous polymucleotide to a predetermined endogenous DNA target sequence in a eukarvotic 10 cell with high efficiency and with sequence specificity. Exogenous polynucleotides, are localized (or targeted) to one or more predetermined DNA target sequence(s) by homologous pairing in vivo. Such targeted homologous pairing of exogenous polynucleotides to endogenous DNA seguences in vivo may be 15 used: (1) to target chemical substituents in a sequencespecific manner in vivo. (2) to correct or to generate genetic mutations in endogenous DNA sequences by homologous recombination and/or gene conversion. (3) to produce homologously targeted transgenic animals at high efficiency. 20 and (4) in other applications (e.g., targeted drug delivery) based on in vivo homologous pairing. Some embodiments of the invention employ targeted exogenous polynucleotides to correct endogenous mutant gene alleles in human cells; the invention provides methods and compositions for correcting disease 25 alleles involved in producing human genetic diseases, such as inherited genetic diseases (e.g., cystic fibrosis) and neoplasia (e.g., neoplasms induced by somatic mutation of an oncogene or tumor suppressor gene, such as p53, or viral genes associated with diease or neoplasia, such as HBV genes). In one embodiment, at least one exogenous

polynucleotide is targeted to a predetermined endogenous DNA sequence and alters the endogenous DNA sequence, such as a chromosomal DNA sequence, typically by targeted homologous recombination within and/or flanking the predetermined endogenous DNA sequence. Generally, two complementary exogenous polynucleotides are used for targeting an endogenous DNA sequence. Typically, the targeting polynucleotide(s) are introduced simultaneously or contemporaneously with one or more

recombinase species. Alternatively, one or more recombinase species may be produced in vivo by expression of a heterologous expression cassette in a cell containing the preselected target DNA sequence.

It is another object of the invention to provide methods whereby at least one exogenous polynucleotide containing a chemical substituent can be targeted to a predetermined endogenous DNA sequence in a metabolically-active eukaryotic cell, permitting sequence-specific targeting of chemical substituents such as, for example: cross-linking 10 agents, metal chelates (e.g., iron/EDTA chelate for iron catalyzed cleavage), topoisomerases, endonucleases, exonucleases, ligases, phosphodiesterases, photodynamic porphyrins, free-radical generating drugs, quinones, 15 chemotheraneutic drugs (e.g., adriamycin, doxirubicin), intercalating agents, base-modification agents, immunoglobulin chains, oligonuclectides, and other substituents. The methods of the invention can be used to target such a chemical substituent to a predetermined DNA sequence by homologous 20 pairing for various applications, for example: producing sequence-specific strand scission(s), producing sequencespecific chemical modifications (e.g., base methylation, strand cross-linking), producing sequence-specific localization of polypeptides (e.g., topoisomerases, helicases, proteases), 25 producing sequence-specific localization of polynucleotides (e.g., loading sites for replication factors, transcription factors and/or RNA or DNA polymerases), and other applications.

It is another object of the present invention to provide methods for correcting a genetic mutation in an endogenous DNA target sequence, such as a sequence encoding a protein. For example, the invention can be used to correct genetic mutations, such as base substitutions, additions, and/or deletions, by converting a mutant DNA sequence that encodes a non-functional, dysfunctional, and/or truncated polypeptide into a corrected DNA sequence that encodes a functional polypeptide (e.g., has a biological activity such as an enzymatic activity, hormone function, or other biological property). The methods and compositions of the invention may

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11 also be used to correct genetic mutations or dysfunctional

alleles with genetic lesions in non-coding sequences (e.g., promoters, enhancers, silencers, origins of replication, splicing signals). In contradistinction, the invention also 5 can be used to target DNA sequences for inactivating gene expression; a targeting polynucleotide can be employed to make a targeted base substitution, addition, and/or deletion in a structural or regulatory endogenous DNA seguence to alter expression of one or more genes, typically by knocking out at least one allele of a gene (i.e., making a mutant. nonfunctional allele). The invention can also be used to correct disease alleles, such as a human or non-human animal CFTR gene allele associated with cystic fibrosis, by producing . a targeted alteration in the disease allele to correct a disease-causing lesion (e.g., a deletion). It is a further object of the invention to provide methods and compositions for high-efficiency gene targeting of human genetic disease alleles, such as a CFTR allele associated

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with cystic fibrosis or an LDL receptor allele assocaited with 20 familial hypercholesterolemia. In one aspect of the invention, targeting polynucleotides having at least one associated recombinase are targeted to cells in vivo (i.e., in an intact animal) by exploiting the advantages of a receptor-mediated uptake mechanism, such as an asialoglycoprotein receptor-25 mediated uptake process. In this variation, a targeting polynucleotide is associated with a recombinase and a celluptake component which enhances the uptake of the targeting polynucleotide-recombinase into cells of at least one cell type in an intact individual. For example, but not limitation, a 30 cell-uptake component typically consists essentially of: (1) a galactose-terminal (asialo-) glycoprotein (e.g., asialogrosomucoid) capable of being recognized and internalized by specialized receptors (asialoglycoprotein receptors) on hepatocytes in vivo, and (2) a polycation, such as poly-Llysine, which binds to the targeting polynucleotide, usually by electrostatic interaction. Typically, the targeting polynucleotide is coated with recombinase and cell-uptake component simultaneously so that both recombinase and celluptake component bind to the targeting polymucleotide; alternatively, a targeting polymucleotide can be coated with recombinase prior to incubation with a cell-uptake component; alternatively the targeting polymucleotide can be coated with the cell-uptake component and introduced into cells contemporaneously with a separately delivered recombinase (e.g., by targeted liposomes containing one or more recombinase).

The invention also provides methods and compositions 10 for treatment and prophylaxis of genetic diseases of animals, particularly mammals, wherein a recombinase and a targeting polynucleotide are used to produce a targeted sequence modification in a disease allele of an endogenous gene. The invention may also be used to produce targeted sequence 15 modification(s) in a non-human animal, particularly a non-human mammal such as a mouse or rabbit, which create(s) a disease allele in a non-human animal. Sequence-modified non-human animals harboring such a disease allele may provide useful models of human and veterinary disease(s). Alternatively, the 20 methods and compositions of the invention can be used to provide non-human animals having homologous Iv-targeted human disease alleles integrated into a non-human genome; such nonhuman animals may provide useful experimental models of human genetic disease, including neoplastic diseases and 25 atherosclerotic disease.

It is also an object of the invention to provide
methods and compositions for recombinase-enhanced positioning
of a targeting polymoclecitie to a homologous sequence in an
endogenous chromosome to form a stable multistrand complex, and
thereby alter expression of a predetermined gene sequence by
interfering with transcription of sequence(s) adjacent to the
multistrand complex. Recombinase(s) are used to ensure correct
homologous pairing and formation of a stable multistrand
complex, which may include a double-D loop structure. For
example, a targeting polymuclectide coated with a recombinase
may homologously pair with an endogenous chromosomal sequence
in a structural or regulatory sequence of a gene and form a
stable multistrand complex which may: (1) constitute a

significant physical or chemical obstacle to formation of or procession of an active transcriptional complex comprising at least an RNA polymerase, or (2) after the local chromatin structure so as to alter the transcription rate of gene sequences within about 1 to 500 kilobases of the multistrand complex.

It is another object of the invention to provide
methods and compositions for treating or preventing human and
animal diseases, particularly viral diseases, such as human
lo hepatitis B virus (HBV) hepatitis, by targeting viral gene
sequences with a recombinase-associated targeting
polymucleotide and thereby inactivating said viral gene
sequences and inhibiting viral-induced pathology.

Tt is a further object of the invention to provide some propositions that contain exogenous targeting polymucleotides, complementary pairs of targeting polymucleotides, chemical substituents of such polymucleotides, and recombinase proteins used in the methods of the invention. Such compositions may include cell-uptake components to facilitate intracellular outside of a targeting polymucleotide, especially for <u>ex vivo</u> and in vivo cene therapy, transcenseis, and ene modification.

BRIEF DESCRIPTION OF THE DRAWINGS

Figs. 1A-1D. Homologous targeting of rec2-coated

stromosome 1 alpha-satellite polymucleotides in living cell
nuclei. The homologously targeted biotinylated polymucleotides
were visualized by addition of FITC-widin followed by washing
to remove unbound FITC. Signals were visualized using a Zeiss
Confocal Lawer Scanning Microscope (CLSM-10) with 488 nm argon
localized FITC-DNA signals in cell nucleus. Fig 1B - enhanced
image of FITC-DNA signals in cell nucleus. Fig 1B - enhanced
image of FITC-DNA signals in cell nucleus. Fig 1C - image of
FITC-DNA signals overlaid on phase image of nucleus. Fig 1D phase image of center of cell nucleus showing nucleoli. Note:
11 images except lower right were photographed at same focus
level (focus unchanged between these photos).

Figs. 2A-2C. Homologous targeting of recA-coated

chromosome 1 alpha-matellite polymorlectides in living cell nuclei. Fig. 2h - fluorescent image of FITC-DNA signals in cell nucleus. Fig. 25 - enhanced image of FITC-DNA signal in cell nucleus. Fig. 2C - everlay of FITC-DNA signals on phase 5 image of nucleus.

Figs. 3A-1B. Decondensed DNA from a targeted human chromosome I in a living cell nucleus displaying repeated alpha-satellite DNA sequences as visualized by FITC-DNA labeling; Fig. 3A - short exposure; Fig. 3B- long exposure.

10 Fig. 4. Human p53 tumor suppressor gene targeting in living HED-2 cell nuclei.

Figs. 5A and 5B. FITC-localization of read-coated polynucleotides targeted to human chromosome 1 alpha-satellite sequences in a living cell nucleus. Fig. 5A - image of 15 enhanced FITC-signals. Fig. 5B - overlay of FITC-signals on phase contrast insec of cell nucleus.

Fig. 6. Map of mammalian expression lacZ mutant plasmid pMC1lacXpa.

Fig. 7. Map of mammalian expression lacZ wild-type
20 plasmid pMC1lacpa.

Fig. 8. Multiple cloning site of plasmid pIBI30. Fig. 9. PCR products and primers from lacZ gene

Fig. 9. FCR products and primers from lacz gene sequence.

Fig. 10A. Southern hybridization analysis of the

25 687-bp fragment amplified from genomic DNA. Electrophoretic migration of a 687-bp DNA fragment generated with primers CP1 and CP5 from genomic DNA of EOTES-Go-cells which were capillary needle-microin/sected with the 491-mucleotide fragment in the presence of reck (lane 2) or transfected as a protein-DNA-lipid complex where the 491-mucleotide fragments were conted with reck (+; lane 3). The control DNA was amplified from montransfected EOTES-Go-cultures (lane 1).

Fig. 108 Autoradiographic analysis of DNA transferred to Gene Screen Plus filters and hybridized with a ¹²P-labeled oligonucleotide specific for normal exon 10 sequences in the region of the *AFSOS mutation. Cells transfected by microinjection or protein-lipid-DNA complexes both were positive for homologous targeting, whereas control cells were not.

CF1/N.

Fig. 11A. Analysis of DNA from cells electroporated or transfected with DNA encapsulated in a protein-lipid complex. Aliela-specific PCR amplification of the 687/684 bp 5 DNA fragment amplified in the first round with primer CF1 and oligo N (N) or oligo AF (AF). Ethidium bromide-stained 300 bp DNA fragment separated by electrophoresis in a 1% agarose gel. The DNA in each lane is as follows: lane 1, 100-bp marker DNA; lane 2, control 168DE140-cell DNA amplified with the CF1/N primers; lane 3, nontransfected XCFTE290-cell DNA amplified with CF1/AF primers; lane 4, nontransfected XCFTE290-cell DNA amplified with CF1/AF primers; lane 5, DNA from XCFTE290-cells electroporated with CF1/AF primers; lane 6, DNA from XCFTE290-cells transfected with CF1/AF primers; lane 6, DNA from XCFTE290-cells transfected with reck-coated 491-nuclectide fragment sensepsulated in a protein-lipid complex and amplified with

Fig. 11B Autoradiographic analysis of the DNA in Fig.
11A transferred to Gene Screen Plus filters and hybridized with
20 Sep-labeled cligo N probe. Samples in lanes 1-5 for the
autoradiographic analysis are equivalent to lanes 2-6 in Fig.
11A.

Fig. 12. PCR analysis of ECTTE290-genomic DNA reconstructed with the addition of 2 x 10⁵ copies of rechcoated 491-nucleotide fregments per microgram of genomic DNA. This number of DNA fregments represents the total number of DNA copies microinjected into cells and tests whether the 491nucleotide fregment can act as a primer for the 687/684-bp fragment amplification. DNA was amplified as described in Fig. 10. When the second round of amplification was conducted with CP1 and the oligo N primers (lane 2), the 300-bp DNA band was not detected when aliquots of the amplification reaction were separated electrophoretically. Amplification of the ECTPE290491 bp DNA fragment with the CP1/oligo AF primer pair produced a 299-bb DNA product (lane 1). Marker DNA is in lane 3.

Fig. 13 graphically portrays the relationship between the number of targeting polymucleotides introduced into cells and the frequency of ones correction using targeting

polynucleotides coated with reck or without reck. The indicated plasmids were microinjected into living cells at differing concentrations. The average number of plasmid molecules injected per cell was calculated from the 5 concentration of plasmid in the fixed volume (approximately 0.5 picolitre) of DNA solution injected per cell. Symbology used is: X = wild-type plasmid; 0 = mutant plasmid pMC1lacXpa + wild-type 276-mer DNA coated with RecA protein; □ = mutant plasmid pMC1lacXpa + wild-type 276-mer DNA without RecA protein.

DEFINITIONS

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Unless defined otherwise, all technical and scientific terms used herein have the same meaning as commonly 15 understood by one of ordinary skill in the art to which this invention belongs. Although any methods and materials similar or equivalent to those described herein can be used in the practice or testing of the present invention, the preferred . methods and materials are described. For purposes of the 20 present invention, the following terms are defined below.

As used herein, the twenty conventional amino acids and their abbreviations follow conventional usage (Immunology -A Synthesis, 2nd Edition, E.S. Golub and D.R. Green, Eds., Sinauer Associates, Sunderland, Massachusetts (1991), which is 25 incorporated herein by reference).

As used herein, the terms "predetermined endogenous DNA sequence" and "predetermined target sequence" refer to polynucleotide sequences contained in a eukaryotic cell. Such sequences include, for example, chromosomal sequences (e.g., 30 structural genes, promoters, enhancers, recombinatorial hotspots, repeat sequences, integrated proviral sequences), episomal sequences (e.g., replicable plasmids or viral replication intermediates), chloroplast and mitochondrial DNA sequences. By "predetermined" it is meant that the target 35 sequence may be selected at the discretion of the practitioner on the basis of known or predicted sequence information, and is not constrained to specific sites recognized by certain sitespecific recombinases (e.q., FLP recombinase or CRE

recombinase). In some embodiments, the predetermined

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endogenous sequences.

endogenous DNA target sequence will be other than a naturally occurring germline DNA sequence (e.g., a transgene, parasitic, or mycoplasmal or viral sequence). An exogenous polynucleotide is a polynucleotide which is transferred into a eukaryotic cell but which has not been replicated in that host cell; for example, a virus genome polynucleotide that enters a cell by fusion of a virion to the cell is an exogenous polynucleotide, however, replicated copies of the viral polynucleotide subsequently made in the infected cell are endogenous sequences (and may, for example, become integrated into a cell chromosome). Similarly, transgenes which are microinjected or transfected into a cell are exogenous polynucleotides, however integrated and replicated copies of the transgene(s) are

a polynucleotide sequence is homologous (i.e., is identical, not strictly evolutionarily related) to all or a portion of a. reference polynucleotide sequence, or that a polypeptide sequence is identical to a reference polypeptide sequence. In contradistinction, the term "complementary to" is used herein to mean that the complementary sequence is homologous to all or a portion of a reference polynucleotide sequence. For illustration, the nucleotide sequence "TATAC" corresponds to a 25 reference sequence "TATAC" and is complementary to a reference sequence "GTATA".

The term "corresponds to" is used herein to mean that

The following terms are used to describe the sequence relationships between two or more polynucleotides: "reference

sequence", "comparison window", "sequence identity", "percentage of sequence identity", and "substantial identity". A "reference sequence" is a defined sequence used as a basis for a sequence comparision: a reference sequence may be a subset of a larger sequence, for example, as a segment of a full-length cDNA or gene sequence given in a sequence listing, such as a human cystic fibrosis gene (Riordan et al. (1989) Science 245: 1066; Rommens et al. (1989) Science 245: 1059), or may comprise a complete full-length cDNA or gene sequence. Generally, a reference sequence is at least 18 nucleotides in

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length, typically at least 25 nucleotides in length, frequently at least 50 nucleotides in length. Since two polynucleotides may each (1) comprise a sequence (i.e., a portion of the complete polynucleotide sequence) that is similar between the 5 two polynucleotides, and (2) may further comprise a sequence that is divergent between the two polynucleotides, sequence comparisons between two (or more) polynucleotides are typically performed by comparing sequences of the two polynucleotides over a "comparison window" to identify and compare local 10 regions of sequence similarity. A "comparison window", as used herein, refers to a conceptual segment of at least 18 contiquous nucleotide positions wherein a polynucleotide sequence may be compared to a reference sequence of at least 18 contiguous nucleotides and wherein the portion of the 15 polynucleotide sequence in the comparison window may comprise additions or deletions (i.e., gaps) of 20 percent or less as compared to the reference sequence (which does not comprise additions or deletions) for optimal alignment of the two sequences. Optimal alignment of sequences for aligning a 20 comparison window may be conducted by the local homology algorithm of Smith and Waterman (1981) Adv. Appl. Math. 2: 482, by the homology alignment algorithm of Needleman and Wunsch (1970) J. Nol. Biol. 48: 443, by the search for similarity method of Pearson and Lipman (1988) Proc. Natl. Acad. Sci. 25 (U.S.A.) 85: 2444, by computerized implementations of these algorithms (GAP, BESTFIT, FASTA, and TFASTA in the Wisconsin Genetics Software Package, Genetics Computer Group, 575 Science Dr., Madison, WI), or by inspection, and the best alignment (i.e., resulting in the highest percentage of homology over the 30 comparison window) generated by the various methods is selected. The term "sequence identity" means that two polynucleotide seguences are identical (i.e., on a nucleotideby-nucleotide basis) over the window of comparison. The term "percentage of sequence identity" is calculated by comparing 35 two optimally aligned sequences over the window of comparison. determining the number of positions at which the identical nucleic acid base (e.g., A, T, C, G, U, or I) occurs in both sequences to yield the number of matched positions, dividing

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the number of matched positions by the total number of positions in the window of comparision (i.e., the window size), and multiplying the result by 100 to yield the percentage of sequence identity. The term "substantial identity" may be used 5 to denote a characteristic of a polynucleotide sequence, wherein the polynucleotide comprises a sequence that has at least 80 percent sequence identity compared to a reference sequence over a comparison window of at least 18 nucleotide positions, wherein the percentage of sequence identity is calculated by comparing the reference sequence to the polynucleotide sequence which may include deletions or additions which total 20 percent or less of the reference sequence over the window of comparison.

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The terms "substantially corresponds to" or 15 "substantial identity" as used herein denotes a characteristic of a nucleic acid sequence, wherein a nucleic acid sequence has at least about 70 percent sequence identity as compared to a reference sequence, typically at least about 85 percent sequence identity, and preferably at least about 95 percent to 20 97 percent sequence identity as compared to a reference sequence, frequently having at least 99 percent sequence identity. The percentage of sequence identity is calculated excluding small deletions or additions which total less than 20 percent of the reference sequence. The reference sequence may 25 be a subset of a larger sequence, such as a portion of a gene or flanking sequence, or a repetitive portion of a chromosome.

long, typically at least about 30 nucleotides long, and preferably at least about 50 to 100 nucleotides long. 30 "Substantially complementary" as used herein refers to a sequence that is complementary to a sequence that substantially corresponds to a reference sequence. In general, targeting efficiency increases with the length of the targeting

However, the reference sequence is at least 18 nucleotides

polynucleotide portion that is substantially complementary to a 35 reference sequence present in the target DNA.

"Specific hybridization" is defined herein as the formation of hybrids between a targeting polynucleotide (e.g., a polynucleotide of the invention which may include

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substitutions, deletion, and/or additions as compared to the predetermined target DNA sequence) and a predetermined target DNA, wherein the targeting polynucleotide preferentially hybridizes to the predetermined target DNA such that, for 5 example, a discrete band can be identified on a Southern blot of DNA prepared from eukaryotic cells that contain the target DNA sequence, and/or a targeting polynucleotide in an intact nucleus localizes to a discrete chromosomal location characteristic of a unique or repetitive sequence. In some 10 instances, a target sequence may be present in more than one target polynucleotide species (e.g., a particular target sequence may occur in multiple members of a gene family or in a known repetitive sequence). It is evident that optimal hybridization conditions will vary depending upon the sequence 15 composition and length(s) of the targeting polynucleotide(s) and target(s), and the experimental method selected by the practitioner. Various quidelines may be used to select appropriate hybridization conditions (see, Maniatis et al., Molecular Cloning: A Laboratory Manual (1989), 2nd Ed., Cold Spring Harbor, N.Y. and Berger and Kimmel, Methods in 20 Enzymology, Volume 152, Guide to Molecular Cloning Techniques (1987), Academic Press, Inc., San Diego, CA., which are incorporated herein by reference. Methods for hybridizing a targeting polynucleotide to a discrete chromosomal location in 25 intact nuclei are provided herein in the Detailed Description. The term "naturally-occurring" as used herein as

applied to an object refers to the fact that an object can be found in mature. For example, a polymnolectide sequence that is present in an organisms (including viruses) that can be isolated from a source in mature and which has not been intentionally modified by man in the laboratory is naturallyoccurring.

A metabolically-active cell is a cell, comprising an intact nucleus, which, when provided nutrients and incubated in an appropriate medium carries out DNA synthesis and ENA for extended periods (e.g., at least 12-24 hours). Such metabolically-active cells are typically differentiated cells incapable of further cell division (although nuclear division

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and chromosomal replication may occur), although stem cells are also metabolically-active cells.

As used herein, the term "disease allele" refers to an allele of a gene which is capable of producing a recognizable disease. A disease allele may be dominant or recessive and may produce disease directly or when present in combination with a specific genetic background or pre-existing pathological condition. A disease allele may be present in the gene pool or may be generated de novo in an individual by somatic mutation. For example and not limitation, disease alleles include: activated oncogenes, a sickle cell anemia allele, a Tay-Sachs allele, a cystic fibrosis allele, a multidrug resistance (MDR-1) allele, a Gaucher's Type I allele, a Lesch-Nyhan allele, a retinoblastoma-susceptibility allele, a Fabry's disease allele, and a Huntington's chorea allele. As used herein, a disease allele encompasses both alleles associated with human diseases and alleles associated with recognized veterinary diseases. For example, the AF508 CFTR .

allele is a human disease allele which is associted with cystic

As used herein, the term "cell-uptake component"

refers to an agent which, when bound, either directly or indirectly, to a targeting polymucleotide, enhances the intracellular uptake of the targeting polymucleotide into at 25 least one cell type (e.g., hepatocytes). A cell-uptake component may include, but is not limited to, the following; a galactose-terminal (asialo-) glycoprotein capable of being internalised into hepatocytes via a hepatocyte asialoglycoprotein receptor, a polycation (e.g., poly-L-1ysine), and/or a protein-lipid complex formed with the targeting polymucleotide. A cell uptake component may include a viral capsid protein, viral coat protein, or viral envelops

protein. Various combinations of the above, as well as alternative cell-uptake components will be apparent to those of

skill in the art and are provided in the published literature.

DETAILED DESCRIPTION

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Generally, the nomenclature used hereafter and the

laboratory procedures in cell culture, molecular genetics, and nucleic acid chemistry and hybridization described below are those well known and commonly employed in the art. Standard techniques are used for recombinant nucleic acid methods, polynucleotide synthesis, cell culture, and transgenesis. Generally enzymatic reactions, oligonucleotide synthesis, oligonucleotide modification, and purification steps are performed according to the manufacturer's specifications. The techniques and procedures are generally performed according to 10 conventional methods in the art and various general references which are provided throughout this document. The procedures therein are believed to be well known in the art and are provided for the convenience of the reader. All the information contained therein is incorporated herein by 15 reference.

Transgenic mice are derived according to Hogan, et al., "Manipulating the Mouse Embryo: A Laboratory Manual", Cold Spring Harbor Laboratory (1988) which is incorporated herein by reference.

Embryonic stem cells are manipulated according to published procedures (Teratocarcinomas and embryonic stem cells: a practical approach, E.J. Robertson, ed., IRL Press, Washington, D.C., 1987; Zjilstra et al., Nature 342:435-438 (1989); and Schwartzberg et al., Science 246:799-803 (1989), 25 each of which is incorporated herein by reference).

Oligonucleotides can be synthesized on an Applied Bio Systems oligonucleotide synthesizer according to specifications provided by the manufacturer.

30 Targeting Polynucleotides

Targeting polynucleotides may be produced by chemical synthesis of oligonucleotides, nick-translation of a doublestranded DNA template, polymerase chain-reaction amplification of a sequence (or ligase chain reaction amplification), purification of prokaryotic or eukaryotic cloning vectors harboring a sequence of interest (e.g., a cloned cDNA or genomic clone, or portion thereof) such as plasmids, phagemids, YACs, cosmids, bacteriophage DNA, other viral DNA or

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replication intermediates, or purified restriction fragments thereof, as well as other sources of single and double-stranded polynucleotides having a desired nucleotide sequence. Targeting polynucleotides are generally ssDNA or dsDNA, most

5 preferably dsDNA.

Targeting polynucleotides are generally at least about 50 to 100 nucleotides long, preferably at least about 250 to 500 nucleotides long, more preferably at least about 1000 to 2000 nucleotides long, or longer; however, as the length of a 10 targeting polynucleotide increases beyond about 20,000 to 50,000 nucleotides, the efficiency of transferring an intact targeting polynucleotide into the cell decreases. The length of homology may be selected at the discretion of the practitioner on the basis of the sequence composition and 15 complexity of the predetermined endogenous target DNA sequence(s) and quidance provided in the art, which generally indicates that 1.3 to 6.8 kilobase segments of homology are preferred (Hasty et al. (1991) Molec. Cell. Biol. 11: 5586; Shulman et al. (1990) Molec. Cell. Biol. 10: 4466, which are 20 incorporated herein by reference). Targeting polynucleotides have at least one sequence that substantially corresponds to, or is substantially complementary to, a predetermined endogenous DNA sequence (i.e., a DNA sequence of a polynucleotide located in a eukaryotic cell, such as a 25 chromosomal, mitochondrial, chloroplast, viral, episomal, or mycoplasmal polynucleotide). Such targeting polynucleotide sequences serve as templates for homologous pairing with the predetermined endogenous sequence(s), and are also referred to herein as homology clamps. In targeting polynucleotides, such 30 homology clamps are typically located at or near the 5' or 3' end, preferably homology clamps are located at each end of the polynucleotide (Berinstein et al. (1992) Molec. Cell. Biol. 12: 360, which is incorporated herein by reference). Without wishing to be bound by any particular theory, it is believed 35 that the addition of recombinases permits efficient gene targeting with targeting polynucleotides having short (i.e., about 50 to 500 basepair long) segments of homology, as well as with targeting polynucleotides having longer segments of

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homology.

The formation of heteroduplex joints is not a stringent process; genetic evidence supports the view that the classical phenomena of meiotic gene conversion and aberrant 5 meiotic segregation result in part from the inclusion of mismatched base pairs in heteroduplex joints, and the subsequent correction of some of these mismatched base pairs before replication. Observations on recA protein have provided information on parameters that affect the discrimination of 10 relatedness from perfect or near-perfect homology and that affect the inclusion of mismatched base pairs in heteroduplex joints. The ability of recA protein to drive strand exchange past all single base-pair mismatches and to form extensively mismatched joints in superhelical DNA reflect its role in 15 recombination and gene conversion. This error-prone process may also be related to its role in mutagenesis. RecA-mediated pairing reactions involving DNA of \$X174 and G4, which are about 70 percent homologous, have yielded homologous recombinants (Cunningham et al. (1981) Cell 24: 213), although 20 recA preferentially forms homologous joints between highly homologous sequences, and is implicated as mediating a homology search process between an invading DNA strand and a recipient DNA strand, producing relatively stable heteroduplexes at regions of high homology.

Therefore, is is preferred that targeting polymoleculeds of the invention have hosology clamps that are highly homologous to the predetermined target endogenous DNA sequence(s), preferably not containing terminally located missatches, and most preferably stopequin. Typically, targeting open content of the preferable that least one homology clamp that is at least about 25 to 15 mulectices long, and it is preferable that homology clamps are at least about 50 to 100 nucleotides long, and nore preferably at least about 50 to 100 nucleotides long, although the degree of sequence homology on nucleotides long, and nore preferably at least about 100-500 nucleotides long, and and preferable the degree of sequence and the base composition of the targeted sequence will determine the optimal and minimal clamp lengths (e.g., G-C rich sequences are typically more thereodymanically stable and will generally

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require shorter clamp length). Therefore, both homology clamp length and the degree of sequence homology can only be determined with reference to a particular predetermined sequence, but homology clamps generally must be about at least 5 25-50 nucleotides long (Rubnitz and Subramani (1985) Mol. Cell. Biol. 5: 659: Avares et al. (1986) Proc. Natl. Acad. Sci. (U.S.A.) 83: 5199: Campbell et al. (1989) The New Biologist 1: 223) and must also substantially correspond or be substantially complementary to a predetermined target sequence. Preferably, a homology clamp is at least about 50 nucleotides long and is identical to or complementary to a predetermined target sequence. Without wishing to be bound by a particular theory. it is believed that the addition of recombinases to a targeting polynucleotide enhances the efficiency of homologous recombination between homologous, nonisogenic sequences (e.g., between an exon 2 sequence of a albumin gene of a Balb/c mouse and a homologous albumin gene exon 2 seguence of a C57/BL6 mouse), as well as between isogenic sequences. The invention is preferably practiced with a complementary pair of targeting polynucleotides, usually of equal length, which are simultaneously or contemporaneously introduced into a eukaryotic cell harboring a predetermined endogenous target sequence, generally with at least one recombinase protein (e.g., recA). Under most circumstances, it is preferred that the targeting polynucleotides are incubated with recA or other recombinase prior to introduction into a eukaryotic cell, so that the recombinase protein(s) may be "loaded" onto the targeting polynucleotide(s). Incubation conditions for such recombinase loading are described infra, 30 and also in U.S.S.N. 07/755,462, filed 4 September 1991; U.S.S.N. 07/910,791, filed 9 July 1992; and U.S.S.N. 07/520,321, filed 7 May 1990, each of which is incorporated herein by reference. A targeting polynucleotide may contain a sequence that enhances the loading process of a recombinase, for example a recA loading sequence is the recombinogenic nucleation sequence poly-[d(A-C)], and its complement, poly-[d(G-T)]. The duplex sequence $poly[d(A-C) \cdot d(G-T)]_n$, where n is from 5 to 25, is a middle repetitive element in eukaryotic DNA.

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The invention may also be practiced with individual targeting polymucleotides which do not comprise part of a complementary pair. In each case, a targeting polymucleotide is introduced into a eukaryotic cell simultaneously or contemporaneously with a recombinase protein, typically in the form of a conted targeting polymucleotide (i.e., a polymucleotide preincubated with recombinase wherein the recombinase is noncovalently bound to the polymucleotide).

A targeting polynucleotide used in a method of the invention typically is a single-stranded nucleic acid, usually 10 a DNA strand, or derived by denaturation of a duplex DNA, which is complementary to one (or both) strand(s) of the target duplex nucleic acid. The homology clamp sequence preferably contains at least 90-95% sequence homology with the target sequence, to insure sequence-specific targeting of the 15 targeting polynucleotide to the endogenous DNA target. Usually, it is preferable that the homology clamps are isogenic with the target polynucleotide. The single-stranded targeting polynucleotide is typically about 50-600 bases long, although a shorter or longer polynucleotide may also be employed. 20 Alternatively, the targeting polynucleotide may be prepared in single-stranded form by oligonucleotide synthesis methods, which may first require, especially with larger targeting polynucleotides, formation of subfragments of the targeting polynucleotide, typically followed by splicing of the 25 subfragments together, typically by enzymatic ligation.

Recombinase Proteins

Recombinases are proteins that, when included with an exogenous targeting polymucleotide, provides a measurable increase in the recombination frequency and/or localization frequency between the targeting polymucleotide and an endogenous predeteration DNR sequence. In the present invention, recombinase refers to a family of Rech-like recombination proteins all having essentially all or most of the size functions, particularly: (i) the recombinase protein's ability to properly bind to and position targeting polymucleotides on their homologous targets and (ii) the

· ability of recombinase protein/targeting polynucleotide complexes to efficiently find and bind to complementary endogenous sequences. The best characterized recA protein is from E. coli. in addition to the wild-type protein a number of 5 mutant reck-like proteins have been identified (e.g., reck803). Further, many organisms have recA-like recombinases with strand-transfer activities (e.g., Pugisawa et al., (1985) Nucl. Acids Res. 13: 7473; Hsieh et al., (1986) Cell 44: 885; Hsieh et al., (1989) J. Biol. Chem. 264: 5089; Fishel et al., (1988) Proc. Natl. Acad. Sci. USA 85: 3683; Cassuto et al., (1987) Mol. Gen. Genet. 208: 10; Ganea et al., (1987) Mol. Cell Biol. 7: 3124; Moore et al., (1990) J. Biol. Chem. 19: 11108; Keene et al., (1984) Nucl. Acids Res. 12: 3057; Kimiec, (1984) Cold Spring Harbor Symp. 48:675; Kimeic. (1986) Cell 44: 545; 15 Kolodner et al., (1987) Proc. Natl. Acad. Sci. USA 84 :5560; Sugino et al., (1985) Proc. Natl. Acad. Sci. USA 85: 3683; Halbrook et al., (1989) J. Biol. Chem. 264: 21403; Eisen et al., (1988) Proc. Natl. Acad. Sci. USA 85: 7481: McCarthy et . al., (1988) Proc. Natl. Acad. Sci. USA 85: 5854; Lowenhaupt et 20 al., (1989) J. Biol. Chem. 264: 20568, which are incorporated herein by reference. Examples of such recombinase proteins include, for example but not limitation: recA, recAsO3, uvsX, and other recA mutants and recA-like recombinases (Roca, A.I. (1990) Crit. Rev. Biochem. Molec. Biol. 25: 415), sepl 25 (Kolodner et al. (1987) Proc. Natl. Acad. Sci. (U.S.A.) 84: 5560; Tishkoff et al. Molec. Cell. Biol. 11: 2593), RuvC (Dunderdale et al. (1991) Nature 354: 506), DST2, KEM1, XRN1 (Dykstra et al. (1991) Molec. Cell. Biol. 11: 2583), STPa/DST1 (Clark et al. (1991) Molec. Cell. Biol. 11: 2576), HPP-1 (Moore 30 et al. (1991) Proc. Natl. Acad. Sci. (U.S.A.) 88: 9067), other eukaryotic recombinases (Bishop et al. (1992) Cell 69: 439; Shinohara et al. (1992) Cell 69: 457); incorporated herein by reference. RecA may be purified from E. coli strains, such as E. coli strains JC12772 and JC15369 (available from A.J. Clark and M. Madiraju, University of California-Berkeley). These strains contain the reck coding sequences on a "runaway" replicating plasmid vector present at a high copy numbers per cell. The recA803 protein is a high-activity mutant of wildtype reck. The art teaches several examples of recombinase proteins, for example, from Drosophila, yeast, plant, human, and non-tunan mammalian cells, including proteins with biological properties similar to reck (i.e., reck-like 5 recombinases).

Recombinase protein(s) (prokaryotic or eukaryotic) may be exogenously administered to a eukaryotic cell simultaneously or contemporaneously (i.e., within about a few hours) with the targeting polynucleotide(s). Such 10 administration is typically done by microinjection, although electroporation, lipofection, and other transfection methods known in the art may also be used. Alternatively, recombinase proteins may be produced in vivo from a heterologous expression cassette in a transfected cell or transgenic cell, such as a 15 transgenic totipotent embryonal stem cell (e.g., a murine ES cell such as AB-1) used to generate a transgenic non-human animal line or a pluripotent hematopoietic stem cell for reconstituting all or part of the hematopoietic stem cell population of an individual, or a terminally differentiated 20 cell type, such as a fibroblast, epithelial cell (e.g., pneumocyte), or muscle cell. Conveniently, a heterologous expression cassette includes a modulatable promoter, such as an ecdysone-inducible promoter-enhancer combination, an estrogeninduced promoter-enhancer combination, a CMV promoter-enhancer, 25 an insulin gene promoter, or other cell-type specific, developmental stage-specific, hormone-inducible, or other modulatable promoter construct so that expression of at least one species of recombinase protein from the cassette can by modulated for transiently producing recombinase(s) in vivo 30 simultaneous or contemporaneous with introduction of a targeting polynucleotide into the cell. When a hormoneinducible promoter-enhancer combination is used, the cell must have the required hormone receptor present, either naturally or as a consequence of expression a co-transfected expression 35 vector encoding such receptor.

For making transgenic non-human animals (which include homologously targeted non-human animals) embryonal stem cells (ES cells) are preferred. Murine ES cells, such as AB-1

line grown on mitotically inactive SNLT6/7 cell feeder layers (McMahon and Bradley, Cell 52:1073-1085 (1990)) essentially as described (Robartson, E.J. (1987) in Teratocartinomas and Suntromic Stem Cells: A Practical Approach E.J. Robertson, ed. 5 (Oxford: IRL Press), p. 71-112) may be used for homologous gene targeting. Other suitable ES lines include, but are not limited to, the E14 line (Hooper et al. (1987) Nature 326: 292-295), the D3 line (Doetschman et al. (1987) in Embryol. Exp. Morph. £2: 27-45), and the CCE line (Robertson et al. (1986) in Sature 322: 445-445). The success of generating a mouse line from ES cells bearing a specific targeted mutation depends on the pluripotence of the ES cells (i.e., their ability, once injected into a host blastocyst. to participate in embryogenesis and contribute to the gers cells of the resulting sanimal).

The pluripotence of any given ES cell line can vary with time in culture and the care with which it has been handled. The only definitive assay for pluripotence is to detarmine whether the specific population of ES cells to be used for targeting can give rise to chimeras capable of germline transmission of the ES genome. For this reason, prior to gene targeting, a portion of the parental population of AB-1 cells is injected into CSTB/(S) lastrocysts to ascertain whether the cells are capable of generating chimeric nice with extensive ES cell contribution and whether the majority of these chimeras can transmit the ES genome to propeny.

The vectors containing the DNA segments of interest can be transferred into the host cell by well-known methods, depending on the type of cellular host. For example, although calcium phosphate treatment, electroporation, lipofection, blolistics or viral-based transfection also may be used. Other methods used to transferor manusclian cells include the use of Polybrene, protoplast fusion, and others (sea, smarrally, Sambrook et al. Molecular Cloning: A Laboratory Manual; Ad ed., 1989, Cold Spring Harbor Laboratory Press, cold Spring Harbor; N.Y., which is incorporated herein by reference). Direct injection of DNA and/or recombinase-costed

targeting polynucleotides into target cells, such as skeletal or muscle cells also may be used (Wolff et al. (1990) <u>Science</u> 247: 1465, which is incorporated herein by reference).

Reck protein is typically obtained from bacterial strains that overproduce the protein: wild-type F. coli reck protein and nutant reckeds protein may be purified from such strains. Alternatively, reck protein can also be purchased from, for example, Pharmacia (Picactaway, NJ).

Reca protein forms a nucleoprotein filament when it coats a single-stranded DNA. In this nucleoprotein filament, one monomer of reca protein is bound to about 1 nucleotides. This property of recA to cost single-stranded DNA is essentially sequence independent, although particular sequences favor initial loading of recA onto a polymncleotide (e.g., nucleotion sequences). The nucleoprotein filament(s) can be formed on essentially any DNA molecule and can be formed in cells (e.g., nammalian cells), forming complexes with both sincle-stranded and double-stranded DNA.

20 Recombinase Coating of Targeting Polynucleotides

The conditions used to cost targeting polymuclectides with recA protein and ATPyS have been described in commonly assigned U.S.S.N. 07/910,791, filled 9 July 1992; U.S.S.N. 07/755,462, filled 4 September 1991; and U.S.S.N. 07/250,321, of filed 7 May 1990, each incorporated herein by reference. Targeting polymuclectides can be coated using GTPyS, mixes of ATPyS with rATP and/or dATP, or dATP or TATP alone Or in the presence of an rATP generating system (Bochringer Mamhaim). Various mixtures of GTPyS, ATPyS, ATP, ADP, ADTP and/or rATP 30 may be used, particularly preferred are mixes of ATPyS and ATP or ATTP's and ATP.

Reck protein coating of targeting polymoclectides is typically carried out as described in W093/17267 and W093/05178. Briefly, the targeting polymoclectide, whether 35 double-stranded or single-stranded, is denatured by heating in an aqueous solution at 95-100°C for if the ninutes, then placed in an ice bath for 20 seconds to about one minute followed by centrifugation at 0°C for approximately 20 sec, before use WO 93/22443 PCT/US93/03868

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When denatured targeting polynucleotides are not placed in a freezer at -00°C they are usually immediately added to standard reach coating reaction buffer containing ATP75, at room temperature, and to this is added the reck protein.

5 Alternatively, recA protein may be included with the buffer components and ATP/S before the polynucleotides are added. RecA coating of targeting polynucleotide(s) is

initiated by innubating polymnolectide-recA mixtures at 37°C for 10-15 min. RecA protain concentration tested during reaction with polymuclectide varies depending upon polymuclectide size and the amount of added polymuclectide, and the ratio of recA molecule:nuclectide preferably ranges between about 3:1 and 1:3. When single-stranded polymuclectides are recA coated independently of their homologous polymuclectide strange, the mix and unconcentrations of ATP-8 and recA.

respectively, can be reduced to one-half those used with double-stranded targeting polymocleotides (i.e. recA and ATPyS concentration ratios are usually kept constant at a specific concentration of individual polymocleotide strand, depending on whether a single- or double-stranded polymocleotide is used).

RecA protein coating of targeting polynucleotides is

normally carried out in a standard 1X RecA coating reaction buffer. 10% RecA reaction buffer (i.e., 10% AC buffer) consists of: 100 mM Tris acctate (pH 7.5 at 37°C), 20 nM magnesium acctate, 500 mM sodium acctate, 10 nM DTT, and 50% glycerol). All of the targeting polynuclaotides, whether double-stranded or single-stranded, typically are denatured before use by heating to 95-100°C for five minutes, placed on ice for up to one minute, and subjected to centriugation (10,000 rpm) at 0°C for approximately 20 seconds (e.g., in a Tomy centrifuge). Denatured targeting polynucleotides usually are added immediately to room temperature RecA coating reaction buffer

A reaction mixture typically contains the following components: (i) 2.4 mM ATPyS; and (ii) between 1-100 mg/μ l of targeting polynucleotide. To this mixture is added about 1-20 μ l of reach protein per 10-100 μ l of reaction mixture, usually

mixed with ATPyS and diluted with double-distilled H2O, as

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at about 5.2-11.0 mg/ml (purchased from Pharmacia or purified), and is rapidly added and mixed. The final reaction volume for Reac coating of targeting polymucleotide is usually in the range of about 10-500 µl. Reac coating of targeting

5 polynucleotide is usually initiated by incubating targeting polynucleotide-RecA mixtures at 37°C for about 10-15 min. RecA protein concentrations in coating reactions

varies depending upon targeting polynuclectide size and the amount of added targeting polynuclectide: reck protein o concentrations are typically in the range of 5 to 50 pM. When single-stranded targeting polynuclectides are coated with reck, independently of their complementary strands, the concentrations of ATPS and reck protein may optionally be reduced to about one-half of the concentrations used with 5 double-stranded targeting polynuclectides of the same length; that is, the reck protein and ATPS concentration ratios are generally kept constant for a given concentration of individual polynuclectide strands.

The coating of targeting polynucleotides with rech 20 protein can be evaluated in a number of ways. First, protein binding to DNA can be examined using band-shift gel assays (McEntee et al., (1981) J. Biol. Chem. 256:8835). Labeled polynucleotides can be coated with reck protein in the presence of ATPyS and the products of the coating reactions may be separated by agarose gel electrophoresis. Following incubation of recA protein with denatured duplex DNAs the recA protein effectively coats single-stranded targeting polynucleotides derived from denaturing a duplex DNA. As the ratio of recA protein monomers to nucleotides in the targeting polynucleotide 30 increases from 0, 1:27, 1:2.7 to 3.7:1 for I21-mer and 0, 1:22, 1:2.2 to 4.5:1 for 159-mer, targeting polynucleotide's electrophoretic mobility decreases, i.e., is retarded, due to reck-binding to the targeting polynucleotide. Retardation of the coated polynucleotide's mobility reflects the saturation of 35 targeting polynucleotide with reck protein. An excess of reck monomers to DNA nucleotides is required for efficient recA coating of short targeting polynucleotides (Leahy et al.,

(1986) J. Biol. Chem. 261:6954).

A second method for evaluating protein binding to DNA is in the use of nitrocellulose fiber binding assays (Leahy et al., (1986) J. Biol. Chem. 261:6954; Woodbury, et al., (1983) Biochemistry 22(20):4730-4737. The nitrocellulose filter 5 binding method is particularly useful in determining the dissociation-rates for protein: DNA complexes using labeled DNA. In the filter binding assay, DNA:protein complexes are retained on a filter while free DNA passes through the filter. This assay method is more quantitative for dissociation-rate determinations because the separation of DNA: protein complexes from free targeting polynucleotide is very rapid.

A targeting polynucleotide of the invention may

Cell-Uptake Components

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15 optionally be conjugated, typically by noncovalent binding, to a cell-uptake component. Various methods have been described in the art for targeting DNA to specific cell types. A targeting polynucleotide of the invention can be conjugated to essentially any of several cell-uptake components known in the 20 art. For targeting to hepatocytes, a targeting polynucleotide can be conjugated to an asialogrosomucoid (ASOR)-poly-L-lysine conjugate by methods described in the art and incorporated herein by reference (Wu GY and Wu CH (1987) J. Biol. Chem. 262: 4429; Wu GY and Wu CH (1988) Biochemistry 27: 887; Wu GY and Wu 25 CH (1988) J. Biol. Chem. 263: 14621; Wu GY and Wu CH (1992) J. Biol. Chem. 267: 12436; Wu et al. (1991) J. Biol. Chem. 266: 14338; and Wilson et al. (1992) J. Biol. Chem. 267: 963, W092/06180; W092/05250; and W091/17761 which are incorporated herein by reference).

by incubating the targeting polynucleotide with at least one lipid species and at least one protein species to form proteinlipid-polynucleotide complexes consisting essentially of the targeting polynucleotide and the lipid-protein cell-uptake 35 component. Lipid vesicles made according to Felgner (WO91/17424, incorporated herein by reference) and/or cationic lipidization (WO91/16024, incorporated herein by reference) or other forms for polynucleotide administration (EP 465,529,

Alternatively, a cell-uptake component may be formed

incorporated herein by reference) may also be employed as celluptake components. Viral capsid proteins, cost proteins, and envelope proteins (e.g., those promoting membrane fusion; hemagglutinin). Examples of viral glycoproteins which attach to cell surface receptors and lead to internalization and/or membrane fusion include the gB, gC, gD, gZ, gH, and gI virion glycoproteins of HSP-1, and gpl20 of HLY-1.

Typically, a targeting polynucleotide of the invention is coated with at least one recombinase and is conjugated to a cell-uptake component, and the resulting cell 10 targeting complex is contacted with a target cell under uptake conditions (e.g., physiological conditions) so that the targeting polynucleotide and the recombinase(s) are internalized in the target cell. A targeting polynucleotide 15 may be contacted simultaneously or sequentially with a celluptake component and also with a recombinase; preferably the targeting polynucleotide is contacted first with a recombinase, or with a mixture comprising both a cell-uptake component and a recombinase under conditions whereby, on average, at least 20 about one molecule of recombinase is noncovalently attached per targeting polynucleotide molecule and at least about one celluptake component also is noncovalently attached. Most preferably, coating of both recombinase and cell-uptake component saturates essentially all of the available binding 25 sites on the targeting polynucleotide. A targeting polynucleotide may be preferentially coated with a cell-uptake component so that the resultant targeting complex comprises, on a molar basis, more cell-uptake component than recombinase(5). Alternatively, a targeting polynucleotide may be preferentially coated with recombinase(s) so that the resultant targeting complex comprises, on a molar basis, more recombinase(s) than cell-uptake component.

cell-uptake components are included with recombinasecoated targeting polymucleotides of the invention to enhance

55 the uptake of the recombinase-coated targeting
polymucleotide(s) into cells, particularly for in yivo gene
targeting applications, such as gene therapy to treat genetic
diseases, including neoplasis, and targeted homologous

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recombination to treat viral infections wherein a viral sequence (e.g., an integrated hepatitis B virus (HBV) genome or genome fragment) may be targeted by homologous sequence targeting and inactivated. Alternatively, a targeting

5 polynucleotide may be coated with the cell-uptake component and targeted to cells with a contemporaneous or simultaneous administration of a recombinase (e.g., liposomes or immunoliposomes containing a recombinase, a viral-based vector encoding and expressing a recombinase).

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Several disease states may be amenable to treatment or prophylaxis by targeted alteration of heptocytes in vivo by homologous gene targeting. For example and not for limitation. the following diseases, among others not listed, are expected to be amenable to targeted gene therapy; hepatocellular 15 carcinoma, HBV infection, familial hypercholesterolemia (LDL

receptor defect), alcohol sensitivity (alcohol dehydrogenase and/or aldehyde dehydrogenase insufficiency), hepatoblastoma, Wilson's disease, congenital hepatic porphyrias, and inherited disorders of hepatic metabolism. Where targeting of hepatic 20 cells in vivo is desired, a cell-uptake component consisting essentially of an asialoglycoprotein-poly-L-lysine conjugate is preferred. The targeting complexes of the invention which may

significantly increased targeting efficiency produced by 25 association of a targeting polynucleotide with a recombinase which, when combined with a cell-targeting method such as that of W092/05250 and/or Wilson et al. (1992) J. Biol. Chem. 267: 963, provide a highly efficient method for performing in vivo homologous sequence targeting in cells, such as hepatocytes.

be used to target hepatocytes in vivo take advantage of the

For many types of in vivo gene therapy to be effective, a significant number of cells must be correctly targeted, with a minimum number of cells having an incorrectly targeted recombination event. To accomplish this objective, the combination of: (1) a targeting polynucleotide(s), (2) a recombinase (to provide enhanced efficiency and specificity of correct homologous sequence targeting), and (3) a cell-uptake component (to provide enhanced cellular uptake of the targeting poynucleotide), provides a means for the efficient and specific

targeting of cells in vivo, making in vivo homologous sequence targeting, and gene therapy, practicable.

Generally, any predetermined endogenous DNA sequence

Targeting of Endogenous DNA Sequences In Vivo

can be altered by homologous recombination (which includes gene conversion) with an exogenous targeting polynucleotide (or complementary pair of targeting polynucleotides) that has at least one homology clamp which substantially corresponds to or is substantially complementary to a predetermined endogenous DNA target sequence and which is introduced with a recombinase (e.g., recA) into a eukaryotic cell having the predetermined endogenous DNA sequence. Typically, a targeting polynucleotide (or complementary polynucleotide pair) has a portion having a sequence that is not present in the preselected endogenous 15 targeted sequence(s) (i.e., a nonhomologous portion) which may be as small as a single mismatched nucleotide or may span up to about several kilobases or more of nonhomologous sequence. Generally, such nonhomologous portions are flanked on each side by homology clamps, although a single flanking homology clamp may be used. Nonhomologous portions are used to make insertions, deletions, and/or replacements in a predetermined endogenous targeted DNA sequence, and/or to make single or multiple nucleotide substitutions in a predetermined endogenous target DNA sequence so that the resultant recombined sequence (i.e., a targeted recombinant endogenous seguence) incorporates some or all of the sequence information of the nonhomologous portion of the targeting polynucleotide(s). Additions and deletions may be as small as 1 nucleotide or may range up to about 2 to 10 kilobases or more. 30

In one application, a targeting polynucleotide can be used to repair a mutated sequence of a structural gene by replacing it or converting it to a wild-type sequence (e.g., a sequence encoding a protain with a wild-type biological 3 activity). For example, such applications could be used to convert a sickle cell trait allele of a hemoglobin gene to an allele which encodes a hemoglobin molecule that is not susceptible to sickling, by altering the nucleotide sequence WO 93/22443 PCT/US93/03868

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encoding the β-subunit of hemoglobin so that the codon at position 6 of the β subunit is converted Valβ6--Sdluβ6 (Shesely et al. (1991) <u>op.cit.</u>). Other genetic diseases can be corrected, either partially or totally, by replacing, inserting, and/or deleting sequence information in a disease allele using appropriately selected exogenous targeting polynucleotides. For example but not for limitation, the ΔF508 deletion in the human CFFE ones can be corrected by targeted

10 polynucleotide of the invention.

Gene Inactivation

In addition to correcting disease alleles, exogenous targeting polynucleotides can be used to inactivate one or more sense in a cell (or transgenic nonhuman animal).

homologous recombination employing a recA-coated targeting

Once the specific target genes to be modified are selected, their sequences will be scanned for possible disruption sites (convenient restriction sites, for example). Plasmids are engineered to contain an appropriately sized gene 20 sequence with a deletion or insertion in the gene of interest and at least one flanking homology clamp which substantially corresponds or is substantially complementary to an endogenous target DNA sequence. Vectors containing a targeting polynucleotide sequence are typically grown in E. coli and then 25 isolated using standard molecular biology methods, or may be synthesized as oligonucleotides. Direct targeted inactivation which does not require vectors may also be done. When using microinjection procedures it may be preferable to use a transfection technique with linearized sequences containing 30 only modified target gene sequence and without vector or selectable sequences. The modified gene site is such that a homologous recombinant between the exogenous targeting polynucleotide and the endogenous DNA target sequence can be identified by using carefully chosen primers and PCR, followed by analysis to detect if PCR products specific to the desired targeted event are present (Erlich et al., (1991) Science 252: 1643, which is incorporated herein by reference). Several studies have already used PCR to successfully identify and then

clone the desired transfected cell lines (Zimmer and Gruss, (1989) Nature 338:150; Mouellic et al., (1990) Prog. Natl. Acad. Sci. USA 87:4712; Shasely et al., (1991) Prog. Natl.

Acad. Sci. USA 82:4294, which are incorporated herein by
reference). This approach is very effective when the number of
cells receiving exogenous targeting polymucleotide(s) is high

cells receiving exogenous targeting polymenlectide(s) is high (i.e., with microinjection, or with liposomes) and the treated cell populations are allowed to expand to cell groups of approximately 1 x 10⁴ cells (capechi, (1989) <u>Science</u> 10 <u>244</u>:1289). When the target gene is not on a sex chromosome, or

244:1288). When the target gene is not on a sex chromosome, or the cells are derived from a female, both alleles of a gene can be targeted by sequential inactivation (Mortensen et al., (1991) Proc. Natl. Acad. Sci. USA 88:7036).

Homologous Pairing of Targeting Polynucleotides Having Chemical Substituents

Exogenous targeting polynucleotides that have been

modified with appended chemical substituents may be introduced along with recombinase (e.g., recA) into a metabolically active eukaryotic cell to homologously pair with a predetermined 20 endogenous DNA target sequence in the cell. Typically such exogenous targeting polynuclectides are derivatized, and additional chemical substituents are attached, either during or after polynucleotide synthesis, respectively, and are thus localized to a specific endogenous target sequence where they 25 produce an alteration or chemical modification to a local DNA sequence. Preferred attached chemical substituents include: cross-linking agents, psoralen, metal chelates (e.g., iron/EDTA chelate for iron catalyzed cleavage), topoisomerases, 30 endonucleases, exonucleases, ligases, phosphodiesterases, photodynamic porphyrins, chemotherapeutic drugs (e.g., adriamycin, doxirubicin), intercalating agents, basemodification agents, immunoglobulin chains, and oligonucleotides. Iron/EDTA chelates are particularly preferred chemical substituents where local cleavage of a DNA

sequence is desired (Hertzberg et al. (1982) J. Am. Chem. Soc. 104: 313; Hertzberg and Dervan (1984) Biochemistry 23: 3934; Taylor et al. (1984) Tetrahedron 40: 457; Dervan, PB (1986)

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Science 232: 464, which are incorporated herein by reference). Preferred attachment chemistries include: direct linkage, e.g., via an appended reactive amino group (Corev and Schultz (1988) Science 238: 1401, which is incorporated herein by reference) 5 and other direct linkage chemistries, although streptavidin/biotin and digoxigenin/anti-digoxigenin antibody linkage methods may also be used. Methods for linking chemical substitutents are provided in U.S. Patents 5,135,720, 5,093,245, and 5,055,556, which are incorporated herein by reference. Other linkage chemistries may be used at the

The broad scope of this invention is best understood with reference to the following examples, which are not intended to limit the invention in any manner.

EXPERIMENTAL EXAMPLES

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EXAMPLE 1

Homologous Targeting of rech-Coated Chemically-Modified Polynucleotides in Cells

Homologously targeted exogenous targeting polynuclotides specifically target human DNA sequences in intact nuclei of metabolically active cells. RecA-coated complementary exogenous targeting polynucleotides were introduced into metabolically active human cells encapsulated 25 in agarose microbeads and permeabilized to permit entry of DNA/protein complexes using the Jackson-Cook method (Cook, P.R. (1984) EMBO J. 3: 1837; Jackson and Cook (1985) EMBO J. 4: 919; Jackson and Cook (1985) EMBO J. 4: 913; Jackson and Cook (1986) J. Mol. Biol. 192: 65; Jackson et al. (1988) J. Cell. 30 Sci. 90: 365, which are incorporated herein by reference). These experiments were designed to specifically target homologous DNA sequences with recA protein in intact nuclei of metabolically active human HEp-2 cells.

Jackson and Cook previously demonstrated that the nuclear membranes of human or other cells may be permeabilized without loss of metabolic function if the cells are first encapsulated in a gel of agarose microbeads. The agarose

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metabolically active cells.

microbead coat contains the cell constituents and preserves native conformation of chromosocomal DNA, while permitting diffusion of macromolecules into and out of the cell compartment. Wittig et al. (1991) Proc. Natl. Acad. Sci. [18.2 2259, which is incorporated herein by reference, demonstrated that monoclonal antibodies directed spainst left-handed 2-DNA could be diffused into these agarose-embedded cells, and that the antibodies were specifically targeted to chromosomal sequences and conformations. In a similar manner, ow incubated biotim- or FITC-labeled complementary DNA targeting polymuclectides coated with read with agarose-coated cell nuclei and verified the correct homologous targeting of the exogenous targeting polymuclectides to specific predetermined human DNA sequences in cell muclei of

Reck-mediated homologous gene targeting with complementary oligonucleotides in intact human cell nuclei was verified directly by homologous targeting using targeting polynucleotides that were hiotinylated. These were of subsequently labeled with a fluorescent compound to verify homologous pairing at specific locations having the predetermined sequence(s). Reck-coated targeting polynucleotides for human chromosoms 1 pericentrometric alpha-estellite DNA sequences were specifically targeted to that were perseabilitied and suspended in saurose.

In these experiments, reck-coated biotinylated exogenous targeting polymucleotides containing homologous sequences to human chromosome 1 alpha satellite DNA were 30 incubated with human HEP-2 cells. The cells were embedded in agazone, then treated with standard buffers (according to Jackson and Cook, op.cit.) to remove the cytoplasmic membrane and cytoplasm immediately before the addition of targeting polymucleotide coated with reak protein.

The experiments were performed with the following results.

First, in order to test protocols to be used in nuclear encapsulation, freshly trypsinized growing human HEp-2 WO 93/22443 PCT/US93/03868

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tumor cells were suspended in complete DMEM encapsulated in a mixture of agarose (2.5%, Fisher-Bioteck) and complete DMEM media adapting the protocols of Nilsson et al., 1983, so that the final agarose concentration was 0.5% (4 volumes cells in 5 suspension with 1 volume 2.5% agarose), and the final cell concentration range was approximately 2.4×10^7 to 8×10^5 . The encapsulated cells in agarose "beads" were placed in petri dishes to which DMEM complete media was added and were allowed to grow for 24 hr in an incubator at 37°C , 7% CO2. At 24 hr,

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the cells were clearly growing and multiplying and thus were An aliquot of agarose containing cells (in beads in DMEM medium) was treated to remove the cytoplasmic membrane and cytoplasm by addition of ice-cold sterile PBS, New Buffer (Jackson et al. (1988) op.cit.: 130 mM KCl, 10 mM Na2HPO4, 1 mM

alive and metabolically active.

of untreated control cells.

MgCl2, 1 mM Na2ATP, and 1 mM dithithreitol, pH 7.4), New Buffer with 0.5 % Triton-X 100, New Buffer with 0.2% BSA, then was centrifuged at low speed using protocols developed by Jackson and Cook, 1985 and 1986 op.cit.; Wittig et al. (1989) J. Cell. Biol. 108: 755: Wittig et al. (1991) op.cit.) who have 20 shown that this treatment allows the nuclear membrane to remain morphologically intact. The nuclei are metabolically active as shown by a DNA synthesis rate of 85 to 90% compared with that

demonstrated by their morphology and exclusion of 0.4% trypan blue. Nuclei in agarose were returned to the humidified CO. incubator at 37°C for 24 hr and remained metabolically active. 30 We observed that filtered, sterile mineral oil used in the emulsification process was difficult to remove entirely and interfered with the microscopic visualization of suspended nuclei. Therefore, the cell-agarose suspension process was simplified. In subsequent experiments cells were gently vortexed with melted (39°C) agarose, then the agarose-cell mixture was sterilely minced before New Buffer treatments. This simpler process, eliminating the oil step, makes it easier to visualize the cells and chromosomes at the completion of

Cytoplasm was effectively removed by the above treatment, and the encapsulated nuclei were intact as

reactions.

After mincing of the agar and New Buffer treatments of the cells, the above protocols were used to homologously target endogenous DNA sequences in encapsulated nuclei as follows: 16.5 µl rech-coated (or non-rech-coated control) nick-translated DNA (labeled with biotin-14-dATP) targeting polynucleotide was prepared and bound under standard native recA protocols (see WO91/17267 and WO93/05178). Minced agarose fragments were centrifuged and New Buffer supernatant removed. 10 The fragments were resuspended in 1 X AC buffer in a 1.5-ml Eppendorf tube, then centrifuged for removal of the buffer (leaving an estimated 50 to 75 gl of buffer), and prepared targeting polynucleotide was mixed with the fragments of agarose-containing nuclei. Reactions were incubated in a 37°C water bath for 2 to 4 hr, then washed, incubated in standard 15 preblock solution, then in preblock supplement with 10 µg/ml FITC-avidin (Vector, DCS grade), and again washed. Experimental results were analyzed by placing a minute amount. of a reaction with 3 to 4 gl antifade on a slide with a slide cover and viewing it by using the Zeiss CLSM-10 confocal laser scanning microscope (CLSM). Completed reactions were also stored refrigerated for later examination.

In the first in vivo experiment, metabolically active HEp-2 cells suspended in 1 × PBS were encapsulated in agarose by gentle vortexing, treated using New Buffer protocols, then 25 incubated for 3 hr 15 min with 100 ng of reck-coated targeting polynucleotide specific for Chromosome 1 alpha-satellite DNA biotinylated with bio-14-dATP by nick translation (BRL, Nick Translation System) using pUC 1.77 plasmid DNA (a 1.77 kb long ECORI fragment of human DNA in the vector pUC9; Cooke et al. 30 (1979) Nucleic Acids Res. 6: 3177: Emmerich et al. (1989) Exp. Cell. Res. 181: 126). We observed specific targeting by the alpha-satellite targeting polynucleotide to pericentromeric chromosome 1 targets in intact nuclei of metabolically active 35 cells. The number and types of signals were essentially identical to those using the same targeting polynucleotide with methanol (or ethanol)-fixed HEp-2 cell targets in suspension. Figures 1A-1D and 2A-2C show specific targeting signals in

20 sequences.

metabolically active cells from this experiment. In the second in vivo experiment, cells suspended in incomplete DMEM media instead of 1 x PBS were encapsulated in agarose and treated with 62.5 ng of the same targeting polynucleotide used in the first experiment described above and 62.5 ng of a freshly biotinylated targeting polynucleotide prepared under the same protocols. In this experiment, the minced agarose fragments were not resuspended in 1 x AC buffer before addition of targeting polynucleotide and some nuclei disintegrated, especially with subsequent centrifugation. The results show that in the nuclei that remained intact, the targeting polynucleotides coated with recA specifically targeted predetermined human DNA targets. In contrast, targeting polynuclectides in reactions without recA did not 15 correctly target the predetermined human DNA sequences. When the targeted DNA (generated from the recA-coated targeting polynucleotides) was decondensed from the nuclei, the alpha-satellite repeat sequences showed precise and evenly spaced signals along the "string" of the alphoid satellite DNA

Thus, the rech-coated targeting polynucleotides were targeted to the repetitive alpha satellite sequences of Chromosome 1. This result showed DNA targeting in intact nuclei to specific human Chromosome 1 sequences. An example of 25 the experimentally extended DNA with specific alpha-satellite signals appears in Figures 3A and 3B.

In the third experiment, cells were suspended in 1 x PBS or in incomplete DMEM media before vortexing with agarose and were tested using 62.5 ng of targeting polynucleotide in 30 reactions with and without rech protein. In addition, the reactions were divided in half and washed and FITC-avidin treated in either buffer adjusted to pH 7 or pH 7.4. Cells were incubated with the recA coated targeting polynucleotide for 3 hr 25 min. Live nuclei treated with targeting 35 polynucleotide alone without recA showed no signals. In the rech-treated reactions, relatively weaker signals were observed in nuclei incubated in 1 × PBS, whereas very strong specific signals were present in nuclei that had been incubated in

incomplete DMEM. There was clearly significantly more signal present in nuclei that were washed and treated with FITC-avidin at pH 7.4 compared with nuclei incubated at pH 7.0. Figures 5A and 5B show nuclei that were treated with recA coated targeting polynucleotides and incubated at both pH 7.4 and 7.5

In a fourth experiment, HEp-2 cells were embedded in agarces prepared with 1x PBS, New Buffer treated, then treated with 100 neg of hotchinglated targeting polymoulocitide complementary to Chromosome 1 alpha-satellite DNA. Controls in 10 this experiment also included reactions without recA protein and additional control reactions supplemented with an identical amount of BSA protein to replace the recA protein. Additionally, cells were also embedded in agarces prepared with 1x AC Duffer. Examples of specific targeting to endogenous

target sequences were 'recorded.

In a fourth experiment, we directly determined if the embedded nuclei (under the conditions used above) were metabolically active. The nuclei in agarose were incubated with bio-21-cVP in complete medium, then incubated for 2 days in the hundifield OO, atmosphere. Bio-21-cVP was incorporated in RNA. After 2 days at 37 °C, the cells were incubated with FITC-etreptavidin and examined. FITC was specifically associated with nucleoil indicative of riboscana RNA biosynthesis, thus directly showing metabolic activity in these businesses the same of the same cells. Similar results were obtained using DNA procursors to measure DNA synthesis. In this experiment it was clear that the majority of nuclei in the FBS agarose reaction had condensed chromosoms. There was nuclear division in a

number of these nuclei also, indicative of full metabolic

30 viability, which was also shown in the AC buffer-treated cells. A fifth experiment was performed using, again, HEp-2 cells embedded in agarose. Final concentration of the cells in agarose was 3.7 x 10⁶/ml. The cells were suspended in 1 x PBS prior to combining with agarose. The final agarose concentration was 0.5s. There were two reactions, one in which recA was used to coat targeting polymucleotide, the second in which recA protein was replaced by BGA at the same protein concentration followed by New Buffer treatments to remove the

oytoplasm. The nuclei in squrose were incubated for 3 hr with targetting polynucleotide, then processed for detection of correctly targeted polynucleotide using the protocols describe previously. FITC-avidin was used to visualise the biotinylated targeting polynucleotide at a concentration of 20 µg/al.

Results showed that cells with the reck-costed complementary targeting polynucleotide displayed specific signals in 25% or more of the intact nuclei. In contrast, the BSA-treated controls did not show any signal.

calls in agarose from this experiment were further incubated at 37°C in the CO2 incubator in complete needium. At 12 hr, these cells were metabolically active. Chromosomes were condensed, and a number of muclei were in the process of dividing. In these experiments, a significant number of the cells incubated with reck-coated complementary targeting polymuclactides showed specific signal, whereas O of the cells incubated with targeting polymuclactide alone showed specific signal.

In summary, rech-coated biotinvlated targeting

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20 polymucleotides for human chromosome 1 alpha-satallite DNA were specifically targeted to human HEp-2 epithelial carcinoma chromosomal DNA in intact cell muclei of metabolically active cells that had been suspended in agarces, then treated with buffers and rech-coaded targeting polymucleotides under 25 suitable reaction conditions (<u>MBUERA</u> and W091/17267 and W093/05738). Specific binding by the reck-coated targeting polymucleotide to chromatin alpha-satellite DNA was observed only in the sparces embedded muclei which were incubated with reck-coated targeting polymucleotides. Control muclei and the coated targeting polymucleotides. Control muclei and the coated targeting polymucleotides in the absence of rech and/or with nonspecific protein exhibited no signal.

Targeting of Human p53 Gene

We performed reck-mediated homologous targeting of 35 biotimylated targeting polymucleotides that were homologous to the human p53 tumor supressor gene, and compared the results to targeting of alpha satellite DMA sequences in human chromosome 1. In these experiments, exponentially growing cells were trypsinised, washed, suspended in incomplete medium and encapsulated in agarose. The agarose was minced into pieces with a razor blade and the encapsulated cells were treated with New Buffer. A sample from each group was removed to verify 5 that nuclei were intact.

Nuclei were washed in 1 x AC buffer and incubated with reck-coated complementary single-stranded DNA oligonucleotides (i.e., exogenous targeting polynucleotides) for 3.5 hours at 37°C. The alpha satellite DNA targeting polynucleotides for chromosome 1 were previously described and were nick-translated with biotinylated deoxyribonucleotides (bio-14-dATP). The p53 tumor suppressor gene polynucleotide was obtained from Oncor (209 Perry Parkway, Gaithersburg, MD 20877) and is a 1.2 kilobase cDNA fragment from a wild-type human p53 gene (Fields and Jang, (1990) Science 242: 1046; 15 Miller et al. (1986) Nature 319: 783; Zakut-Houre et al. (1985) EMBO J. 4: 1251). The 1.2 kilobase human p53 DNA was nicktranslated with biotinylated deoxyribonuclectides and yielded a population of biotinylated targeting polynucleotides having a size range (about 100 to 600 nucleotides) similar to that obtained for the human chromosome 1 alpha satellite targeting polynucleotides. The targeting polynucleotides were separately incubated with encapsulated cells. Following incubation 3 washes of 1.75 x SSC were done, and sampled nuclei were 25 verified as intact after the washing step. After washing, the targeted encapsulated cell nuclei were incubated in preblock and FITC-avidin was added to preblock buffer to a final concentration of 20 mg/ml for 15 minutes in the dark. The targeted encapsulated cell nuclei were washed sequentially in 4 30 x SSC, 4 x SSC with 0.1% Triton X-100,, and then 4 x SSC. Samples of nuclei were again taken and used to verify that the targeted nuclei were metabolically active. Microscopic examination showed that metabolically active cells contained specific FITC-targeting polynucleotide:targeted endogenous sequence complexes (shown in Figure 4). The p53 targeting 35 polynucleotides were specifically targeted to human chromosome 17, the location of the endogenous human p53 gene sequences, indicating specific pairing of a targeting polynucleotide to a

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unique endogenous DNA target sequence. The human chromosome 1 alpha satellite DNA was also specifically targeted to the chromosome 1 pericentromeric satellite sequences.

The experiments validated a highly specific DNA targeting technique for human or other cells as evidenced by homologous sequence targeting techniques in metabolically active cells. The targeting technique employs the unique properties of reck-mediated DNA sequence targeting with single-estranded (complementary) short targeting

10 polymucleotides. Native intact nuclei were incubated with labeled, heat-denatured targeting polymucleotides coated with recA protein. The DNA hybridized to the predetermined targeted homologous sequences. In these experiments, the targeting polymucleotides formed paired complexes with specific gene sequences within metabolically active cell muclei. This in

yivo targeting by reck-mediated homologous targeting polymucleotides shows the targeting specificity and therapeutic potential for this new in yivo methodology. Application of reck or other recombinase-mediated targeting of (complementary) ssDNA or denatured dsDNA targeting polymucleotides to

20 ssDNA or denatured dsDNA targeting polynucleotides to predetermined endogenous DNA targets is important for human gene entry, gene knockout, gene replacement, and gene mutation or correction.

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EXAMPLE 2

Correcting a Nutant Gene to Produce a Functional Gene Product
Romologously targeted complementary DNA
oligonucleotides were used to correct 11 bp insertion mutations
in vector genes and restore vector gene expression and vector
protein function in microinjected mammelian cells.

Experiments were designed to test whether a homologously targeted complementary oligonucleotide targeting polynucleotide of approximately 270-280 bp (276-bp) could correct an 11-bp insertion mutation in the lack gene of a mammalian DNA vector which encoded a nonfunctional β -galactosidase, so that a corrected lack gene encoded and expressed a functional enzyme. Functional enzyme (β -galactosidase) was detected by an X-qal assay that turns

cells expressing a revertant (i.e., corrected) lacZ gens a blue color.

NHH373 cells microinjected with the mutant test vector bearing an 11 basepair insertion in the lac% coding 5 sequence do not produce any detectable functional β -galactosidase enzyme. In contrast, cells microinjected with the wild two test weter do produce functional enzyme.

We obtained the functional lac plasmid NMC11eops for use as a positive control for expression of β -galactosidass. 10 pMC11acNpa is the target test mutant plasmid (shown in Figure 6). It is identical to pMC11acpa (shown in Figure 7) but has a 11-bp Xbarl Inkre insertional swattion. This plasmid does not express β -galactosidase activity in mouse NHH933 cells when introduced by electroperation. It does not produce blue color 15 in the presence of X-GAL indicative of β -galactosidase production following vector microinjection. Regative controls with mock or noninjected cells we also done. Using these controls and NHH933 cells have no detectable background blue stainine.

20 The plasmid pMC1lacpa (8.4 kb) contains the strong polyoma virus promoter of transcription plus ArG placed in front of the lack gene. The polyadenylation signal from 8940 virus was placed in back of the lack gene. The plasmid vector was pIB130 (shown in Figure 8) from IBI (New Haven, CT). The mutant vector pMC1lacpa has a 11-bp insertion at the XbeI site. This mutation consists of the inserted sequence CTCTAGACGCC (see Figure 9).

In several control microinjection experiments using pMCIlaOXpa we consistently failed to detect any blue 30 microinjected cells. In contrast, in various experiments approximately 8 to 134 of the 373 cells injected with pMCIlacpa DNA expressed 6-galactorisiase as evidenced by their blue color. No cells microinjected with injection buffer alone or mock injected were observed as blue.

We synthesized two 20-bp primers for producing a 276-bp PCR product (see Figure 9) from the wild-type lacZ sequence for use as targeting polynucleotides. We chose this fragment to span the 11 bp insertion mutation as a WO 93/22443 PCT/US93/03868

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nonhonologous sequence. The 276-bp DNA oligonucleotide was separated by gel electrophoresis and electroeluted from agarose, ethanol precipitated, and its concentration determined by absorbance at 260 mm. The fragment was 5' end-labeled with ³²P and specifically D-looped with the pMC1lacxpa or pMC1lacxpa plasmid DNA using recA as shown by agarose gel electrophoresis.

Experiments were designed to test for 8-galactoside production in cells microinjected with MC11acXpa vectors with targeting polymucleotide-target complexes using complementary 276-bp oligonucleotide targeting polymucleotide treated with reck. The 276-mer targeting polymucleotides in 1 XTE buffer were denatured by heating at 100°C for 5 min and immediately quenched in an ice bath for 1 min. The DNA solution was collected at 4°C by centrifugation. Reck-mediated targeting

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15 polynucleotide reactions containing a final volume of 10 µl were assembled using 1.0 µl lox Ac buffer, 1.5 µl 16 mM ATPyS, 3.8 µl 40 Hg, 0.1 2 µl reck protein solution (1) mg/µl), and 2.5 µl of a 30 µg/ml stock of heat-denatured 276-bp targeting polynucleotide. The reck protein was allowed to coat the DNA of or 10 min at 37°C. Next. 1.0 al of 10x AC buffer. 1.0 µl of

for 10 min at 37°C. Next, 1.0 pl of 10% AC Buffer, 1.0 pl of 0.2 M magnesium socates, 1.3 pl of pMCINLOXPA (1.0 mg/µl), and 6.7 pl of dd H₂O was added to a final volume of 20 pl. Control reactions were performed without added rech protein.

NHST37 cells were capillary needle microinfected with

25 targeting polymucleotide-target DNA mixtures loaded in glass pipettes freehly pulled into microneedles using a Sutter instruments microprocessor controlled apparatus. An ECET Eppendorf microinjection pump and computerized micromanipulator were used for computer-assisted microinjection using an Olympus 30 INT-2 inverted microscope. Cells were carefully microinjected under controlled pressure and time. NHHTO cells injected with pMCLlacpa showed approximately 90 of the injected cells were blue. None (00) of the cells injected with pMCLlacXpa DNA in reactions containing the 276 bp oligonucleotide but without reach protein showed a blue color. In marked contrast, approximately 10 of the cells microinjected with the rech-mediated 776-bp targeting polymucleotide targeted to the pMCLlacXpa target bybrid were blue. Thus, these measurements

indicate that the mutant pMCLIacXpa gene can be targeted and corrected by the 276-bp oligomuclectide, which has been targeted with rech-coated targeting polymoclectides. In summary, these measurements show that the 11 bp Xba I insertion mutation can be corrected with the rech-mediated targeted corrected in yivo, but not with the 276-bp oligomuclectide alone. Note that the in situ identification of 373 cells expressing β-palactosidase was performed following incubation with X-pal (5-brono-4-chloro-3-indoly1-β- galactopyranoside) (Sigma), as described by Fischer et al. (1988) Natures 122: 853; Price et al. (1987) Proc. Natl. Acad. Sci. (U.S.Al. 84: 156; Lin and Chee (1989) Biotechniques 2: 576.

Table 1 shows the results of an experiment demonstrating the effect of reck coating of targeting polymoclectides on efficiency of gene correction. The combination of the 276-by targeting polymoclectide and reck (Sample 5) produced a significant frequency of correction of the mutant β-galactosidase gene (3.6%) as compared to samples without reck.

TABLE 1

CORRECTION OF AN INSERTION MUTATION IN THE

8-GALACTOSIDASE GENE OF EURARYOTIC EXPRESSION VECTOR

PMCLIACKDA WITH ROCA-COLATED NORMAL 275-MER DNA²

25	Sample	Injected Plasmid, 276-mer DNA and Reca Protein	Number of Injected Surviving Cells ^b	Number of Surviving Cells Scoring Blue	Surviving Cells Scoring Blue (%)	Level of Signifi- cance
	1	PSV-β-gal-276- mer-RecA	168	21	12.5	
30 -	2	pMC1lacpa-276- mer-Recă	98	9	9.2	
	3	pMC11acXpa-276- mer-RecA	173	. 0	0	
	4 .	pMC11acXpa+276- mer-Reck	103	. 0	0	

_						
Γ	5	pMCllacXpa+276-	168	6°	3.6	<0.05 ^d
1		mer+RecA				

* NII 372 cells were needle microsisected with tive types of plasmids of plasmids with a wide-type f-plantonises gone (Smillage or PSV-)-plasmid six the wide-type f-plantonises gone (Smillage or PSV-)-plantonises (PSV-) plantonises and the second with a motority of the plantonises of the plantonist of the plantonisty as cally the plantonises are cally the plantonises of the plantonisty as cally the plantonisty of the plantonisty in cells of the plantonisty of the plantonisty in cells of the plantonisty of the plantonisty in cells of the plantonists of the plantonisty in cells of the plantonists of the plantonists

needle microinjected with the wild-type plasmids is shown for comparison.

- b On average, about 50% of the total microinjected cells survived.
- 20 ° Five cells were dark blue and one was light blue.
- 4 The numbers of surviving cells scoring blue in the mutant plaumid Rechtrated and non-Bechtrated samples (3,4 and 5) were compared with fourfold of tests. The frequency of corrected cells in the Rach-treated. sample (Sample 5.6 cores the control of the cont
- Courtoid of tesses. The frequency of corrected cells in the secon-transact sample (Sample 5) four of 168) is significantly higher than that of cells in Sample 5 is significantly higher than that of cells in Sample 5 is significantly higher than that of Sample 4 at the 5 significance level (r = -2.75 r_{ad}). The frequency of corrected cells in the Rech-treated Sample 5 is significantly higher than that of Sample 3 at the 14 significance level (r = 6.25 r_{ad}). When samples 3 and 4 are
- 30 at the 14 significance level (x* = 6.28x*0₆₀). When Samples 3 and 4 are combined and compared with Sample 5, the frequency of corrected cells in the Rech-treated Sample 5 is significantly higher than that of the combined sample at the 0.18 significance level (x* = 9.999x*0₆₀₀₁). These are statistically significant data.
- Fig. 13 shows the relationship between the number of plasmids microlinjected per cell and the correction efficiency, and shows the efficiency increase in gene correction which occurs with the Reck-coated targeting polymoclectide as compared to the equivalent polymoclectide which is not coated with Reck.

EXAMPLE 3

Correcting a Human CFTR Disease Allele Homologously targeted complementary DNA

- 45 oligonucleotides were used to correct a naturally occurring 3 by deletion mutation in a human CFTR allele and restore expression of a functional CFTR protein in targeted mammalian cells.
- A major goal of cystic fibrosis (CF) gene therapy is the correction of mutant portions of the CF transmembrane conductance regulator (CFFR) gene by replacement with wild-type

complex.

1227).

DNA sequences to restare the normal CFTR protein and ion transport function. Targeting polynucleotides that were coated with reak protein were introduced into transformed CF airway epithelial cells, homozygous for both alleles AFSOS CFTR gene 5 mutation, by either intranuclear microinjection, electroporation, or by transfection with a protein-DNA-lipid

Isolation and characterization of the CFTR gene (Rommens et al. (1989) Science 245: 1059; Riordan et al. (1989) Science 245: 1066, incorporated herein by reference) has been crucial for understanding the biochemical mechanism(s) underlying CF pathology. The most common mutation associated with CF, a 3-base-pair, in-frame deletion eliminating a phenylalanine at amino acid position 508 (AF508) of CFTR, has 15 been found in about 70% of all CF chromosomes (Kerem et al. (1989) Science 245: 1073; Kerem et al. (1990) Proc. Natl. Acad. Sci. (U.S.A.) 87: 8447). Correction of AF508 and other CFTR DNA mutations lies at the basis of DNA gene therapy for CF disease. Elimination of the cAMP-dependent Cl ion transport 20 defect associated with CFTR gene mutations has been accomplished through the introduction of the transcribed portion of wild-type CFTR cDNA into CF epithelial cells (Rich

et al. (1990) Nature 347: 358; Drumm et al. (1990) Cell 62:

25 An immortalized CF Tracheobronchial epithelial huma cell line, DCFEE300-, is homosygous for the aF968 mutation (Kunzelmann et al. (1992) Nm. J. Respir. Cell. Mol. Biol., in press). These cells are useful as targets for homologous recombination sanalysis, because they contain the same 3 ob haspagir deletion in CFTE allele on all copies of chromosoms 7. Replacement of the aF96 sallele with wild-type CFTE DNA is indicated only when homologous recombination has occurred. The 491 bp region of the CFTE gene spanning exon 11 and containing 3 and 5' flanking intron sequences was selected from sequence 36 data published previously (Eslenski et al. (3991) Genomics 1214, incorporated herein by reference) and used as a targeting polynucleotids. The DNA fragment was PCR amplified in preparative quantities and then denstured for introduction into

cells as recA-coated complementary ssDNA (or dsDNA).

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Exponentially growing cells were transfected by intranuclear microinjection and were propagated on the same petri dishes in which they were microinjected. Cells outside the microinjected area were removed by scraping with a rubber policeman. Exponentially growing cells were typsinized and washed before electroporation. Cells transfected with protein-DNA-lipid

complexes were grown to approximately 70-80% confluence before transfection.

The 491 bp fragment was generated by PCR amplification from the T6/20 plasmid (Rommens et al. (1989) op.cit., incorporated herein by reference) and verified by restriction enzyme mapping and propagated as described previously. After digestion with EcoRI and HindIII, a 860 bp insert was isolated following electrophoresis in 0.8% SeaPlaque agarose gel. The 860 bp fragment contained CFTR exon 10, as well as 5' and 3' intron sequences, as defined by the restriction enzyme cleavage sites (Zielenski et al. (1991) . op.cit.). A 50 ng aliquot of the fragment was amplified by PCR using primers CF1 and CF5 (Table 2) to generate a 491 bp fragment. The conditions for amplification were denaturation, 94° C for 1 min: annealing, 53° C for 30 sec: extension, 72° C for 30 sec with a 4 sec/cycle increase in the extension time for 40 cycles. The fragment size was confirmed by electrophoresis on a 1% agarose gel, then amplified in bulk in DNA. The 491 bp PCR products were extracted with phenol:chloroform:iscamvl alcohol (25:24:1) extraction and

20 separate PCR amplifications, each containing 50 ng of target precipitated with ethanol. DNA precipitates were collected by centrifugation in an Eppendorf microcentrifuge and resuspended at a final concentration of 1 mg/ml. The 491 bo fragment contained exon 10 (193 bp), as well as 5' (163 bp) and 3' (135 bp) flanking intron sequences, as defined by primers CF1 and CF5.

The 491 nucleotide fragments were coated with recA protein using the reaction buffer of Cheng (Cheng, et al. (1988) J. Biol. Chem. 263: 15110, incorporated herein by reference). Typically, the 491 bp DNA fragment (5ug) was

denatured at 95° C for 10 min, then added to a 51 ml of coating buffer containing 200 mg of reck protein, 4.8 mm ATPyS, and 1.7 ml reaction buffer (100 mm Tris-Ac, pH 7.5 at 37° C; 10 mm dithlothreitol; 500 mm NaAc, 20 mm NgAc, 50 percent glycerol) 5 and incubated for 10 min at 37° C. Next, the MgAc concentration was increased to a final concentration of about 22 mm by addition of 7 ml of 200 mm NgAc. Under these conditions, the 491 mucleotide fragment was coated with recA protein at a molar ratio of 3 bases per 1 recA molecule. After 0 coating the fragments were immediately placed on ice at 4° C until transfection (10 min to 1 hr).

Microinjection, when used, was performed with an Eppendorf 5242 microinjection pump fitted to an Eppendorf 5170 ' micromanipulator using borosilicate pipettes (Brunswick, 1.2 15 OD x 1.9ID) fabricated into a microneedle with a Sutter Instruments (P-87) micropipette puller. The micropipettes were filled by capillary force from the opposite side of the needle. Approximately 100 pipettes were used for injecting of 4000 cells. Cells were injected with approximately 1,000-10,000 fragments per cell by intranuclear injection with 120 hPa for 0.1-0.3 s at a volume of 1-10 fl/nucleus. . Microinjected cells were viewed with an Olympus IMT-2 inverted microscope during the injection. The area of the petri dish containing injected cells was marked with 2 to 5 mm diameter rings. Needle 25 microinjection was performed in cells grown on 10 separate 60 mm petri dishes. Cells were injected at room temperature in culture medium after two washes in phosphate buffered saline (PBS). After microinjection, moninjected cells in the culture were removed by scraping. Injected cells were grown at 37° C 30 in a humidified incubator at 7 days and then harvested for DNA and PNA

Electroporation experiments were performed using reck-coasted 491-mer subMn as described above. Approximately 1 x 10⁸ exponentially growing cells were suspended in 400µl of ocating buffer with 5 µg of reck ocated-LNA. The cell suspension was pre-incubated on ice for 10 ml and electroporated at room temperature with 400 V and 400 µF in a EXX 300 electroporator (SEX CORPORATION. SAN BAGEO. CA). After

. electroporation, cells were incubated on ice for an additional 10 min, diluted in Eagle's minimal essential medium (MEM) supplemented with 10% fetal bovine serum (PBS) and 100 µg/ml streptomycin, 100 U/ml penicillin (Cozens et al. (1992) Proc.

5 Natl. Acad. Sci. (U.S.A.) 89: 5171; Gruenert et al. (1988) Proc. Natl. Acad. Sci. (U.S.A.) 85: 5951; Kunzelmann, (1992) op.cit.), and then seeded in T75 flasks. Under these conditions of elecroporation, approximately 30-50% of the cells survive. Cells were cultured for 507 days at 37° C and then harvested for DNA and RNA.

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Protein DNA-lipid complexes (liposomes) were prepared. Briefly, dioleoylphosphatidylethanolamine (PtdEtn, DOPE) was used for preparing liposomes by drying 4 µM solutions of the lipid under nitrogen at room temperature. The lipid film was rehydrated with 4 ml of 30 mM Tris-HCl buffer (pH 9),

then sonicated for 15 minutes under an atmosphere or argon. The protein-DNA complex was prepared in polystyrene tubes by diluting 20 µg of reck-coated 491-base DNA in 30 mM Tris-HCl. (pH 9) buffer. Protein (gramicidin S peptide, GmS; Legendre et 20 al. (1993) Proc. Natl. Acad. Sci. (U.S.A.) 90: 893) was also diluted with 30 mM Tris HCl (pH 9) to a final concentration of 2 mg/ml from a 20 mg/ml stock solution prepared in dimethyl

sulfoxide. The protein, GmS, (40 µg) was added to the DNA and rapidly mixed. Next, 175 µl of the liposome solution (175 nmoles of lipid) were added to the peptide DNA mixture, forming a gramicidin S peptide DNA lipofection complex.

Genomic DNA was isolated and purified from cells as described in Maniatis op.cit. to test for homologous DNA recombination. Cellular DNA was first PCR-amplified with primers CF1 and CF6 (Table 2). CF1 is within the region of homology defined at the 5' end of the 491 bp CFTR fragment CF6 is outside the region of homology at the 3' end of this fragment.

The conditions for the PCR amplification were as follows: CF1/CF6; 684/687 bp fragment; primers, 0.5 µM; DNA, 1-2 µg; denaturation; 94°C for 1 min; annealing; 53°C for 45 s; extension; 72°C for 90 s with a 4-s/cycle increase in extension time for 40 cycles; Mg*2 1.5 mM. DNA fragments were separated

by agarone electrophoresis and visualized by staining with ethidium bromide, then transferred to Gene Screen Plus filters (DuPont). The DNA was then hybridized with the allele-specific normal CTFR ¹²p-and-labeled DNA probe defined by Oligo N as described by Cozens et al. (1992) <u>op.cit.</u>, Nunselmann (1992) <u>op.cit.</u>, incorporated herein by reference. The presence of wild-type (WT) sequences was determined autoradiographically by hybridization with the radiolabeled DNA probe.

Homologous recombination was verified in a second or round of PCR DNA amplification using the 687/684 bp fragment as a DNA template for amplification. The primers used in this allele-specific reaction were CP1 and the oligo N or oligo AF. The size of the DNA fragments was 300 bp (oligo N) or 299 bp (oligo AF).

The conditions for the reaction were as follows: CF1/oligo N/AF; 300/299 bo fragment; primers, 0.5 µM; DNA, 1-2

μg; denaturation, 95°C for 45s; annealing, 51°C for 30s; extension, 72°C for 30 s with a 3-s/cycle increase in extension time for 40 cycles; Mg¹². 1.5 mM. In DNA from transfacted

ECPTEISO-cells, msplified with the CTI/oligo N primers, a PCR product was detected only if the wild-type CPTE sequences were present. Amplification with the CTI/oligo AP gives a PCR DNA product of DNA targets purified from transfected and nontransfected ECPTEISO-cells but not for DNA targets isolated from control normal cells (16HEHGH-0-). The presence of wild-

type CFTR sequences in the amplified DNA fragments was also determined autoradiographically after hybridization with \$2P-

5'-endlabeled oligo N as probe.

Oytoplasmic RNA was isolated and denatured at 95°C
for 2 min, then reverse-transcribed using the DNA polymerase
provided in a PCR NNA Gene Amp kit according to mamufacturer's
instructions (Perkin-Rimer/Octus). First strand CDNA was
amplified by using primer CF17 at the 5' end of exon 9 and the
allele-specific oligo N or oligo aF primers. The length of the
FOR fragments is 322 bp (CF17/Oligo N) and 321 bp (CF17/Oligo
aF).

The conditions for PCR amplification are CF17/oligo N/AF, 322/321 bp fragment; primers, 1 µM; denaturation, 94°C WO 93/22443 PCT/1/S93/03868

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for 1 min; annealing, 51°C for 30s; extension, 72°C for 20s with a 4-s/cycle increase in extension time for 40 cycles; ${\rm Mg^{*2}}$, 0.8 mM. DNA fragments were visualized after

electrophoresis on ethicium bromide-stained 1% agarose gels.

5 In addition to the allele-specific PCR amplification of firststrand cDNA, Southern hybridization was performed as described
above. Pregments were transferred to Gene Screen Plus filters
then hybridized with allele-specific oligo N probe under the
same conditions used for the Southern analysis of the genomic
DNA (Kunzelmann et al. (1992) ggniti; Josens et al. (1992)
gg.citi.) The presence of wild-type CFTR RNA was confirmed by
hybridization and autoratiography of RNA extracted from normal

(16HBE140-) control DNA and in DNA of transfected ECFTE290cells.

Rybridization was performed as described previously (Cozans et al. (1992) <u>op.oft.</u>). DNA fragments were separated by agarose enel electrophoresis. DNA was denatured with 0.4 N

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NaOH and 0.6 M NaCl for 30 min, then washed once with 1.5 M NaCl and 0.5 M Tris-Hcl for 30 min. DNA was transferred to

Gene Screen Plus membrane (MEN-DuPont) by capillary blot, again denatured with 0.4 N NaON for 1 min, and then neutralized with 0.2 M Tris-NCl (pH 7.0). DNA on membranes was prehybridized for 1 h at 37°C in 6 X SSC, 5 X Demhardt's solution, 1% SDS, containing 100 µg/ml of denatured salmon sperm DNA (Sigman).

25 Oligonuclectide probes (oligo N or oligo AF; 10 ng) were ³²7-5'-endlabeled with 20 units of 74 kinase and 40 µci ³²9-γ-ATP for 30 min at 37°C. Unincorporated nucleotides were removed by centrifugation of the reaction mix through a minispin column (Worthington Biochemical Corp., Freehold, NJ). Rybridization
30 Was parformed overnight at 37°C. Weshryanes were washed tytos

was performed overnight at 37°C. Membranes were washed twice for 5 min each time in 2 x SSC at room temperature, twice for 30 min in 2 x SSC, 0.18 SSD at 45°C, and once in 0.1 x SSC for 30 min at room temperature. After washing, hybrids on membranes were analyzed autoradiographically by exposure to x-ray film.

Analysis of ECFTE290- DNA shows replacement of the endogenous mutant (AF508) sequences with the exogenous normal

endogenous mutant (AF508) sequences with the exogenous normal fragment as evidenced by PCR amplification of genomic DNA and

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allele-specific Southern blot hybridization. PCR primers, one inside (CF1), and one outside (CF6) the region of homology (491 bp), were used to test whether the amplified DNA band was possibly due to amplification of any residual DNA fragment 5 remaining in the cell after the transfection or by possible random DNA integration. A 687 bp fragment contains normal CFTR sequences while the 684 bp fragment is generated from AF508 CFTR DNA. To determine whether endogenous AF508 sequences were replaced with exogenous normal CFTR sequences, we analyzed aliquots of the 687 or 684 bp amplification fragments by Southern hybridization using 32P-end-labeled DNA probes specific for the AF508 or wild-type sequences (Table 2). In addition, the 687 bp fragment was PCR amplified by using the CF6 primer and a primer specific for either AF508 (oligo AF) or 15 normal sequences (oligo N). The second round of DNA amplification with the CF1/oligo N or AF primer pair combination yields 300/299 bp fragments, respectively. With the CF1/oligo N primer pair combination, a fragment will be . detected only if the mutant DNA has been replaced by normal sequences. Further confirmation of homologous DNA recombination was tested by allele-specific Southern blot hybridization of the 300/299 bp fragments

Analysis of cytoplasmic RNA to detect normal exon 10 sequences in CFTR mRNA, verify that the homologous DNA 25 recombination was legitimate and that normal CFTR mRNA is expressed in the cytoplasm. To test whether the PCR generated DNA fragments were exclusively CFTR mRNA-derived, primers in exon 9 (CF17) and allele-specific (normal, oligo N or AF508, oligo AF) primers in exon 10. This amplification with primers 30 CF17/N yields a 322 bp normal fragment only if transcription of homologously recombined DNA has occurred. A 321 bp DNA fragment would be generated if the AF508 mutation were present. Furthermore, Southern hybridization analysis with allelespecific 32p-end-labeled probes differentiated between normal 35 and AF508 mutant sequences and were also used to confirm expression of wild-type CFTR mRNA in the cytoplasm.

Homologous recombination between the targeting polynucleotide comprising WT CFTR DNA and AF508 mutant cellular

59 DNA allelic targets was evaluated by analysis of cellular DNA and RNA isolated from transfected and nontransfected ECFTE29ocell cultures. Nuclear genomic DNA and cytoplasmic RNA were isolated 6 days after transfection, CFTR exon 10 sequences were 5 amplified by PCR. Oligonucleotide primers (Table 2) were used to amplify the region of CFTR DNA spanning exon 10. One PCR primer (CF 1) was within the region of homology defined by the 491 bp DNA fragment (sense primer), and the other (CF 6) was outside the homologous region in the 3' intron (antisense primer). This DNA amplification reaction produces a 687 bp fragment with normal human CFTR DNA or a 684 bp fragment if the DNA contains the aF508 mutation, as shown in Fig. 10A. Southern hybridization was carried out on the 687/684 bp DNA fragments generated from amplification of genomic DNA from cell cultures by microinjection or by transfection with the protein-DNA-lipid complex, shown in Fig. 10B. A probe consisting of ³²P-end-labeled oligonucleotide DNA that hybridized only to DNA sequences generated from a normal exon 10 was used. DNA from all microinjected and transfected cells produced specific hybrids as evidenced by autoradiographic hybridization. For cells microinjected with the 491 nucleotide fragment (Fig. 10B. lane 2), the present of normal exon 10 sequences indicated homologous replacement at at least a frequency of ≥ 2.5 x 10-4. This result indicates at least one correctly targeted homologous DNA replacement in about 4000 microinjected nuclei. Other similar experiments using either electroporation or protein-DNA-lipid transfection to transfer the recA-coated 491 nucleotide CFTR DNA fragments also showed homologous recombination with the normal CFTR sequence in transfected CF 30 cells. No hybridization was observed in control nontransfected (or mock-injected ECFTE290- cells). In each cell transfected with normal CFTR DNA, analysis of the genomic DNA in a second round of allele-specific amplification of the 687/684 bp fragments with primers CF1/oligo N (Table 2) clearly showed the

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35 300 bp fragment expected when wild-type CFTR sequences are present, as shown in Fig. 11A. Fragments were detected for control 16HBE140- cells (Fig. 11A, lane 2) and cells transfected with recA-coated DNA (Fig. 11A, lanes 5 and 6). A

299 bp fragment (AF508-specific primer ends one base closer to the CF1 than the oligo N) was detected in DNA from nontransfected ECFTE29g- cells amplified with CF1/oligo AF primers (Fig. 11A, lane 4). No fragment was detected in DNA from nontransfected ECFTE29o- cells reamplified with the CF1/oligo N primers (Fig. 11A, lane 3). Allele-specific Southern blot hybridization of these fragments with the 32P-endlabeled oligo N probe resulted in autoradiographic hybridization signals from control normal and transfected CF 10 cells (Fig. 11B, lames 1, 4, and 5) but not from DNA of nontransfected CF cells amplified with CF1 and oligo-N or -AF (Fig. 11B lames 2 and 3). We tested whether any residual 491 nucleotide DNA fragments which might remain in the cell after 6 days could act as a primer for the PCR reaction, genomic ECFTE290- DNA was incubated with an equivalent number of reckcoated DNA fragments (103-104) introduced by microinjection (Fig. 12). One antisense primer contains the wild-type normal (N) sequence while the other contains the AF508 (AF) mutation. Amplification with the CF1/AF primer combination gives a 299 bp fragments when the AF508 mutation is present. No DNA fragment product was detected when the CF1/N primer combination was used

with control nontransfected ECFTE290- DNA (Fig. 12, lane 2). However, when the CF1/aF primer combination was used for DNA amplification in nontransfected ECFTE290- cells, a DNA product 25 of the expected size (299 bp) was produced (Fig. 12, lane 1). These results indicate that all residual 491 nucleotide DNA fragments which might remain in the cells after 6 days of culture were incapable of competing with the CF1 PCR primers in the PCR amplification of the 687/684 bp fragments.

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5 Olimonuclentide

Table 2
PCR Primers and Oligonucleotides

DNA Semience

-	OTIGORAGIECTIAE	DHA SCIAIL	DNA Sequence
	CF1	8	5'-GCAGAGTACCTGAAACAGGA
	CF5	A	5'-CATTCACAGTAGCTTACCCA
	CF6	A	5'-
	CCACATATCACTATATGO	CATGC	
10	CF17	s	5'-
	GAGGGATTTGGGGAATT	TTTG	
	OLIGO N	A	· 5 '-CACCAAAGATGATATTTTC
	OLIGO AF	A	5 -AACACCAATGATATTTCTT

DMA Strand

15 Notes:

- (1) CF1 and CF5 PCR primers were used to synthesize
- the 491 bp fragment used for the targeting polynucleotide.

 (2) CF1 and CF6 PCR primers were used to amplify the
 687/684 bp CFTR fragment.
- (3) The CF17 primer is located at the 5' end of exon 9 and was used for amplification of first strand cDNA derived from CFFR mRNA.
 - (4) Oligo N and Oligo AF are allele-specific probes and can also be used as allele-specific PCR primers for
- 25 amplifying the 300/299 bp fragments (DNA analysis) and the 322/321 bp fragments (RNA analysis).
- (5) Sense (S) and antisense (A) primers are designated under DNA Strand and indicate the sense of the strand relative to the transcribed direction (i.e., the CFTR 30 mRNA).

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CFTR RNA seguence in the AF508 region of exon 10. Cytoplasmic RNA was isolated from the cells, reverse-transcribed with DNA polymerase and PCR-amplified as first-strand cDNA. This

The corrected CFTR DNA must also be expressed at the

amplification was performed with a PCR primer located in exon 9 (CF17, sense) and CFTR allele-specific PCR primer in exon 10 (oligo N or AF, antisense). The exon 10 primer contains the CF mutation site, and the resulting fragment is 322 bp in normal DNA or 321 bp in DNA containing the AF508 mutation. Amplification of genomic DNA is eliminated by using primers that require amplification across intron/exon boundaries. Amplified cDNA generated from normal control 16HBE140- cells and experimentally transfected cells vielded DNA product fragments with the CF17/oligo N. whereas nontransfected ECFTE290- cells only showed a DNA fragment after amplification with the CF17/oligo AF primers but not with the CF17/oligo N . primers. Cells electroporated with wild-type 491-mer CFTR DNA showed the presence of wild-type CFTR mRNA. In addition, 20 protein-DNA-lipid-transfected ECFTE290- cell cultures also showed the presence of wild-type CFTR mRNA in cells transfected with the recA-coated 491 nucleotide fragment. Southern hybridization of the 322/321 bp cDNA fragments with the 32pend-labeled N oligonucleotide DNA probe showed the specificity of the PCR amplification and produced specific autoradiographic hybridization signals from all cell cultures transfected with reck-coated 491 nucleotide targeting polynucleotide. No autoradiographic hybridization signals were detected in 30 nontransfected ECFTE29o- cells amplified with the CF17/oligo N or oligo AF primers. These analyses verify that the genomic DNA homologously recombined with the WT 491-mer DNA at the AF508 CFTR DNA locus resulting in RNA expressed and transported

This evidence demonstrates that human CFAF508 epithelial cells CFTR DNA can homologously recombine with targeting polynucleotides comprising small fragments of WT CFTR DNA resulting in a corrected genomic CFTR allele, and that a

to the cytoplasm as wild-type CFTR mRNA.

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recA-coated targeting polymucleotide can be used in transfection reactions in cultured human cells, and that cystic fibrosis aF508 mutations can be corrected in genome DNA resulting in the production of normal CFER cytoplasmic MRNA.

Functional analysis of CFTR Cl Transport in mooktransfected and transfected cells was performed by measurement of ³⁶Cl efflux. Following exposure to the cAMP agonist, forskolin, a slight increase in the rate of Cl efflux was detected in the population of transfected cells. A selection protocol was developed to select for small subpopulations of phenotypically normal cells (i.e., transfected cells having functionally normal Cl ion transport) by depleting intracellular Ca² with a Ca² inosphore in a Ca²-Tree buffer and exposing the cells to hypocompolar and then hyperosolar buffers. Defective CP cells that had not undergone functional correction by homologous recombination with a targeting polymolocotide were thus selected sgainst. Measurement of ³⁶Cl efflux in cells enriched by this selection protocol indicated a significant increase in ³⁶Cl following stimulation

Taken together, the data provided indicates that small (e.g., 491-mer) ssDNA fragments can find their genomic homologues when coated with reck protein and efficiently produce homologously targeted intact mammalian cells having a corrected gene sequence. Analysis of CFTR in cytoplasmic RNA and genomic DNA by allele-specific polymerase chain reaction (PCR) amplification and Southern hybridization indicated wild-type CFTR DNA sequences were introduced at the appropriate nuclear genomic DNA locus and was expressed as CFTR MRNA in 10 transfected cell cultures. Thus, in human CF sirway epithelial cells, 491 nuclectide cytoplasmic DNA fragments can target and

20 of cAMP-dependent Cl transport.

AF508 deletion.

Ocrrectly targeted homologous recombination was

detected in one out of one microinjection experiment with rechcoated targeting polymuclectide, two of two different
electroporation experiments with rech-coated targeting
polymuclectide, and one of one limical-MNA-motain commitex

replace the homologous region of CFTR DNA containing a 3 bo

transfection experiment with read-coated targeting polymuclactide. Taken together, these 4 separate experiments strongly indicate that homologous recombination with relatively small read-coated targeting polymuclactides (491-mer GPTR DNR) is feasible for treatment of human quentic diseases, and can be performed successfully by using various methods for delivering the targeting polymuclactides-recombinase complex.

Although the present invention has been described in some detail by way of illustration for purposes of clarity of 10 understanding, it will be apparent that certain changes and modifications may be practiced within the scope of the claims.

SEQUENCE LISTING

(1) GENERAL INFORMATION:

- (i) APPLICANT: Zarling, David A. Sena, Elissa P.
- (ii) TITLE OF INVENTION: In Vivo Homologous Sequence Targeting in Eukaryotic Cells
- (iii) NUMBER OF SEQUENCES: 12
- (iv) CORRESPONDENCE ADDRESS: (A) ADDRESSEE: Townsend and Townsend
 - (A) ADDRESSE: Townsend and Townsend
 (B) STREET: One Market Plaza, Steuart Tower, Suite 2000
 - (C) CITY: San Francisco
 - (D) STATE: California (E) COUNTRY: USA
 - (F) ZIP: 94105
 - (v) COMPUTER READABLE FORM: (A) MEDIUM TYPE: Floppy disk
 - (B) COMPUTER: IBM PC compatible (C) OPERATING SYSTEM: PC-DOS/MS-DOS
 - (D) SOFTWARE: PatentIn Release #1.0, Version #1.25
- (vi) CURRENT APPLICATION DATA:
 - (A) APPLICATION NUMBER: US 07/939,767
 - (B) FILING DATE: 02-SEP-1992 (C) CLASSIFICATION:
- (vii) PRIOR APPLICATION DATA:
 - (A) APPLICATION NUMBER: US 07/873,438
 - (B) FILING DATE: 04-APR-1992
- (viii) ATTORNEY/AGENT INFORMATION:
 - (A) NAME: Smith, William M.
 - (B) REGISTRATION NUMBER: 30,223 (C) REFERENCE/DOCKET NUMBER: 0287A-24-1
 - (ix) TELECOMMUNICATION INFORMATION:
 - (A) TELEPHONE: 415-326-2400
 - (B) TELEFAX: 415-326-2422

(2)	INFO	RMATIC	ON FO	R S	EQ ID	NO:1	.:
	(i)	SEQUE	ENCE	CHAI	RACTE	RIST	ccs
		(A)	LENG	TH:	162	pase	pa

(B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: DNA (genemic) (xi) SEQUENCE DESCRIPTION: SEQ ID NO:1:

ATGATTACGG	ATATCGAATT	AATACGACTC	actataggga	GATCGAATTC	GAGCTCGGTA	60
CCCGGGGATC	CTCTAGAGTC	GACCTGCACC	TGCAGGGGCC	CTCGAGACGC	GIGGCAIGCA	120

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(2)) INFORMATION	FOR	SEQ	ID	NO:2:

CGTAAGTGAA GCGACCCGCA

- (i) SEQUENCE CHARACTERISTICS:
- (A) LENGTH: 20 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: DNA (primer)
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:2:

20

(2)	INFORMATION	FOR	SEO	ID	NO:3:

AGGCCARTCC GCGCCGGATG C

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 21 base pairs
 (B) TYPE: nucleic acid
 (C) STRANDEDNESS: single
 (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: DNA (primer)

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:3:

(2)	INFORMATION	FOR	SEQ	ID	NO:4:

CGATACACCG CATCCGGCGC GGATTGGCCT

(i)	SECUENCE	CHARACTERISTICS:

- EQUENCE CHARACTERISTICS:
 (A) LENGTH: 270 base pairs
 (B) TYPE: nucleic acid
 (C) STRANDEDNESS: single
 (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: DNA (genomic)

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:4:

(/						
CGTAAGTGAA	GCGACCCGCA	TTGACCCTAA	CGCCTGGGTC	GAACGCTGGA	AGGCGGCGGG	60
CCATTACCAG	GCCGAAGCAC	GGTTGTTGCA	GTGCACGGCA	GATACACTTG	CTGATGCGGT	120
GCTGATTACG	ACCECTCACE	CGTGGCAGCA	TCAGGGGAAA	ACCTTATTTA	TCAGCCGGAA	180
AACCTACCGG	ATTGATGGTA	GTGGTCAAAT	GGCGATTACC	GTTGATGTTG	AAGTGGCGAG	240

TATEODERATION		

(i)	SEQUI	ENCE (HA	RACT	ERI	ST	cs:
	(A)	LENG	TH:	281	ba	se	pai
	(B)	TYPE	: n	ucle	ic	ac:	d

- (C) STRANDEDNESS: single (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: DNA (genomic)

(x1) SEQUENCE DESCRIPTION: SEQ ID NO:5: .

				GAACGCTGGA		60
CATTACCAG	GCCGAAGCAC	GGTTGTTGCA	GTGCACGGCA	GATACACTTG	CTGATGCGGT	120
CTGATTACG	ACCGCTCACG	CGCTCTAGAC	GCGTGGCAGC	ATCAGGGGAA	AACCTTATTT	180
ATCAGCCGGA	AAACCTACCG	GATTGATGGT	AGTGGTCAAA	TGGCGATTAC	CGTTGATGTT	240
SAAGTGGCGA	GCGATACACC	GCATCCGGCG	CGGATTGGCC	T		281

(2)	INFORMATION	FOR	SEQ	ID	NO:6:

- (i) SEQUENCE CHARACTERISTICS:
- (A) LENGTH: 11 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: DNA (primer)
- (xi) SEQUENCE DESCRIPTION: SEQ ID No:6:

CTCTAGACGC G

- (2) INFORMATION FOR SEQ ID NO:7:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 20 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: DNA (primer) .
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:7: GCAGAGTACC TGAAACAGGA

- (2) INFORMATION FOR SEQ ID NO:8:
 - (i) SEQUENCE CHARACTERISTICS:

 - (A) LENGTH: 20 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: DNA (primer)
 - (iv) ANTI-SENSE: YES

-

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:8:

CATTCACAGT AGCTTACCCA

- (2) INFORMATION FOR SEQ ID NO:9:
 - (i) SEQUENCE CHARACTERISTICS:

 - (A) LENGTH: 22 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: DNA (primer)
 - (iv) ANTI-SENSE: YES

CCACATATCA CTATATGCAT GC

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:9:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 22 base pairs
 - (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear
- (D) TOPOLOGY: linear (ii) MOLECULE TYPE: DNA (primer)
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:10:

GAGGGATTTG GGGAATTATT TG

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121	TARODMATTON	EUB	SEO	TD	NO:11:

- (i) SEQUENCE CHARACTERISTICS:
 (A) LENGTH: 19 base pairs
 (B) TYPE: nucleic acid
 (C) STRANDENNESS: single
 (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: DNA (primer)
- (iv) ANTI-SENSE: YES
- (x1) SEQUENCE DESCRIPTION: SEQ ID NO:11: CACCAAAGAT GATATITIC

(2)	INFORMATION	FOR	SEQ	ID	NO:12:

- (i) SEQUENCE CHARACTERISTICS:

 - (A) LENGTH: 20 base pairs
 (B) TYPE: nucleic acid
 (C) STRANDEDNESS: single
 (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: DNA (primer)
- (iv) ANTI-SENSE: YES
- (xi) SECUENCE DESCRIPTION: SEC ID NO:12:

AACACCAATG ATATTTTCTT

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CLATMS

- A method for targeting and altering, by homologous recombination, a pre-selected target DNA sequence in a eukaryotic cell to make a targeted sequence modification, said
- 5 eukaryotic cell to make a targeted sequence modi method comprising the steps of:

introducing into at least one eukaryotic cell at least one recombinase and at least one targeting polymucleotide having a homology clame that substantially corresponds to or is sequence; and

identifying a eukaryotic cell having a targeted DNA sequence modification at a preselected target DNA sequence.

- 15 2. A method according to Claim 1, wherein at least two targeting polynucleotides which are substantially complementary to each other are used.
- A method according to Claim 1, wherein said
 recombinase is a species of prokaryotic recombinase.
 - 4. A method according to Claim 3, wherein said prokaryotic recombinase is a species of prokaryotic recA protein.
 - A method according to Claim 4, wherein said reck protein species is E. coli reck.
 - A method according to Claim 1, wherein said targeting polynucleotide is conjugated to a cell-uptake component.
 - A method according to Claim 6, wherein said celluptake component is conjugated to said targeting polynucleotide by noncovalent binding.
 - A method according to Claim 6, wherein the celluptake component comprises an asialoglycoprotein.

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- A method according to Claim 6, wherein the celluptake component comprises a protein-lipid complex.
- 10. A method according to Claim 6, wherein said targeting 5 polynucleotide is conjugated to a cell-uptake component and to a recombinase, forming a cell targeting complex.
 - A method according to Claim 1, wherein said targeting polynuclectide comprises a homology clamp that is complementary to said preselected target DNA sequence.
 - 12. A method according to Claim 11, wherein the targeting polynucleotide consists of a homology clamp.
- 15 13. A method according to Claim 2, wherein a first said targeting polymuclectide comprises a homology clamp that is complementary to said preselected target DNA sequence and a second said targeting polymuclectide comprises a homology clamp that corresponds to said preselected target DNA sequence.
- 14. A method according to Claim 13, wherein said first targeting polynucleotide consists of a homology clamp.
- 15. A method according to Claim 13, wherein the homology 25 clamp of said first targeting polynucleotide and the homology clamp of said second targeting polynucleotide are complementary.
 - 16. A method according to Claim 2, wherein a first said of targeting polynucleotide comprises a homology clamp that is complementary to a preselected target DNA sequence.
 - 17. A method according to Claim 16, wherein a second targeting polymucleotide comprises a homology clamp that is complementary to a sequence of said first targeting polymucleotide.
 - 18. A method according to Claim 17, wherein said second

targeting polynucleotide consists of a sequence that is complementary to the complete sequence of said first polynucleotide.

- 5 19. A method according to Claim 1, wherein the preselected target DNA sequence is unique in a haploid genome of said eukaryotic cell.
- 20. A method according to Claim 19, wherein the preselected target DNA sequence is unique in a diploid genome of said sukaryotic call.
 - A method according to Claim 1, wherein the targeted sequence ordification comprises a deletion of at least one additional nucleatide.
 - 22. A method according to Claim 1, wherein the targeted sequence modification comprises the addition of at least one additional nucleotide.
 - 23. A method according to Claim 22, wherein the targeted sequence modification corrects a human disease allele in a human cell.
 - 5 24. A method according to Claim 23, wherein the human disease allele is a CFTR allele associated with cystic fibrosis.
- 25. A method according to Claim 1 or Claim 6, wherein the 3 recombinase and the targeting polynucleotide are introduced into the eukaryotic cell simultaneously.
- 26. A method according to Claim 25, wherein the recombinase and the targeting polynucleotide are introduced into the eukaryotic cell by a method selected from the group consisting of: microinjection, electroporation, or contacting of the cell with a lipid-protein-targeting polynucleotide complex.

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27. A method according to Claim 1, wherein the targeted sequence modification creates a sequence that encodes a polypeptide having a biological activity.

- 5 28. A method according to Claim 27, wherein the biological activity is an enzymatic activity.
 - 29. A method according to Claim 27 or Claim 28, wherein the targeted sequence modification is in a human cell and encodes a human polypertide.
 - 30. A method according to Claim 29, wherein the targeted sequence modification is in a human oncogene or tumor suppressor gene sequence.
- 31. A method according to Claim 30, wherein the targeted sequence modification is in a human p53 sequence.
- A method according to Claim 29, wherein the targeted
 sequence modification is in a human CFTR allele.
 - 33. A method according to Claim 32, wherein the targeted sequence modification occurs in a human cell.
- 25 34. A method according to Claim 1, wherein the targeting polynuclaotide comprises a homology clamp that is less than 500 nucleotides long.
- 35. A method according to Claim 34, wherein the targeting 30 polynucleotide is less than 500 nucleotides long.
 - 36. A composition for producing a targeted modification of an endogenous DNA sequence, comprising a targeting polynucleotide and a recombinase.
 - 37. A composition according to Claim 36, wherein the targeting polynucleotide is noncovalently bound to said recombinase.

- 38. A composition according to Claim 36, further comprising a cell-uptake component.
- 39. A composition for producing a targeted sequence modification of a human disease allele, comprising a targeting polynucleotide containing a corrected sequence and a recombinase or an expression polynucleotide that encodes and expresses a recombinase.
- 10 40. A composition according to Claim 39, further comprising a cell-uptake component.
 - 41. A composition according to Claim 39 or Claim 40, wherein the human disease allele is a CFTR allele.
 - 42. A kit for therapy, monitoring, or prophylaxis of a genetic disease comprising a recombinase and a targeting polynucleotide.
- 20 43. A kit for therapy, monitoring, or prophylaxis of a genetic disease according to Claim 42, further comprising a cell-uptake component.
 - 44. A method for treating a disease of an animal harboring a disease allele, comprising administering to the animal a composition consisting essentially of:
 - a recombinase or an expression polynucleotide encoding a recombinase; and
- a targeting polynucleotide which produces a sequence 30 modification upon homologous recombination with the disease allele.
 - 45. A method according to Claim 44, wherein the composition further comprises a cell-uptake component.
 - 46. An animal comprising an allele that has a sequence modification according to the method of Claim 44.

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FIG. IA.



FIG. IB.

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FIG. IC.



FIG. ID.



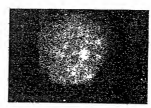


FIG. 2A.

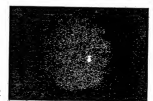


FIG. 2B.



FIG. 2C.

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FIG. 3A.

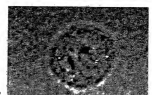


FIG. 3B.



FIG. 4.

SUBSTITUTE SHEET





FIG. 5A.

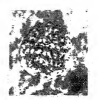


FIG. 5B.

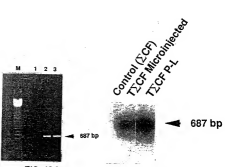


FIG. IOA.

FIG. IOB.

SUBSTITUTE SHEET

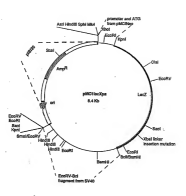


FIGURE 6

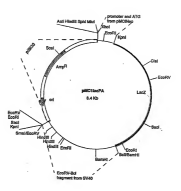


FIGURE 7

Figure 8

pIB1130 multiple cloning site (bases 223-278)

5-ATG ATT ACG CAT ATC GAA TTA ATA CGA CTC ACT ATA CGG AGA TCG AAT TCG AGC TCG GTA CCC GGG GAT CCT CTA GAG TCG ACC TGC AGG GCC CCT CCA GAC GCG TOB CAT GCA AGE TITT CTC CCT TIA GTG AGG GTT AAT TAT AGG CCT AGC TTG-3'

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Figure 9

ATAAAAAACAACTGCTGACGCCTGCGCGATCAGTTCACCCGTGCACCGCTGGATAACG ACATTGGCGTAAGTGAAGCGACCCGATTGACCCTAACGCCTGGGTCGAACGCTGGAAGG TGGCGAGCGATACACCGCATCCGCCCGATTGGCCTGAACTGCCAGGTGG CAGAGGGGTAAACTGGCTCGGATTAGGGCCGCAAGAAAACTATCCCGACCGCCTTACTG COCCOCICATIACCACOCCOAAGCACCOTTOTICCACTCCACCGCAGATACACTTCCTC GCCGGAAAACCTACCGGATTGATGGTAGTGGTCAAATGGCGATTACCGTTTGATGTTGAAG ATGCGGTGCTGATTACGACCGCTCACGCTGGCAGCATCAGGGGAAAACCTTATTATCA PCR B PCR a

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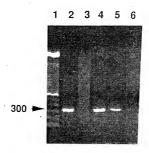


FIGURE 11A

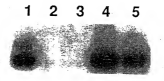


FIGURE 11B

SUBSTITUTE SHEET

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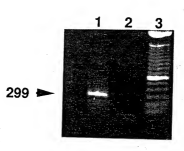


FIGURE 12

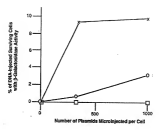


FIGURE 13

PCT/US 93/03868

1 45 10			International Application No	FU1/U3 93/U380
I. CLASSII	ACATION OF SUBJ	ECT MATTER (If several classic)	fication symbols apply, indicate all/4	
According	to International Pater	et Classification (IPC) or to both No		
int.UI.	. 5 C12N15/9 A61K48/0			G01N33/68
O. FIELDS	SEARCHED			
		Misleus	Documentative Searched	
Classificati	an System		Chemification Symbols	
Int.C1.	5	C12N ; A61K A01K	; C12Q ; G0	1N
		Documentation Searche to the Extern that such Doc	of other than Minimum Decumentation names are Indicised in the Fidds Searcher	
III. DOCUM		D TO BE BELEVANT		
Caregory	Contain of De	cusest, a with indicative, where a	ppropriate, of the relevant passages (2	Raissant to Claim No. ¹²
	vol. 17, PRESS, 0 pages 14 B. SAUER recombin sequence genome!	ACIDS RESEARCH no. 1, 11 January XFORD, UK; 77 - 161 AND N. HENDERSON ation at loxP-conts s placed into the; 158, line 9 - page	*Cre-stimulated aining DNA mammalian	1,11,12, 23,28, 29,35, 37,38
	MASHINGT pages 13 S. O'GORI gene act intergra: see page	, 15 March 1991, A/ ON,DC,US; 51 - 1355 MAN ET AL. 'Recombification and site spiton in mammalian of 1351, left column, olumn, line 8	Inase-mediated ecific ecils' line 1 -	1,11,12, 23,28, 29,35,37
			-/	
"A" document of the country of the c	document but publish late set which may throw is cited to establish th or other special reas set referring to an ora setast out published prior to an the priority data of an an a	ments 1 ²⁰ all state of the set which is not it reference it reference of one rather the international leader on printing claim(s) or a publicative date of another on (as specified on (as specified all disclosure, urs, exhibition or this laborational filling date but blined	The formular published efter it or formular published efter of the confidence of the	it the chalcast invention mast he considered to it the chalcast invention at inventive step when the or more other such docu- tivises to a person skilled
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	ENTS CONSIDERED TO BE BELEVANT (CONTINUED FROM THE SECOND SHEET)	
Category o	Citation of Document, with indication, where appropriate, of the relevant passages	Relevant to Claim No
x	J. BACTERIOL. vol. 172, no. 2, February 1990, AM.SOC. MICROBIOL., BALTIMORE, US;	1,11,12, 23,28, 29,35,
	pages 510 - 518 H. MATSUZAKI ET AL. "Chromosome engeneering in Saccharomyces cerevisiae by using a site-specific recombination system of a yeast plasmid	37,38
.	see page 510, left column, line 1 - right column, line 18	
'	J. BIOL. CHEM. vol. 267, no. 2, 15 January 1992, AM. SOC. BIOL. CHEM. INC. US; pages 963 - 967	6-8,10
	J.M. WILSON ET AL. 'Hepatocyte-directed gene transfer in vivo leads to transfent improvement of hypercholesteromia in low density lipoprotein receptor-deficient	
	rabbits' cited in the application see page 965, right column, line 1 - page 967, right column, line 12	8
	NATURE vol. 350, 21 March 1991, MACMILLAM vol. 350, 21 March 1991, MACMILLAM vol. 350, 21 March 1991, MACMILLAM vol. 350, 245 vo	1,3-5, 11,12, 26-30,35
	see page 342, right column, line 1 - page 244, left column, line 9; figures 1-4	-
	WO,A,9 117 267 (SRI INTERNATIONAL) 14 November 1991 cited in the application see page 4, line 5 - line 34; claims 1-18	1,3-8, 10-12, 26-30,35
,х	MO,A,9 208 791 (THE UNITED STATES OF AMERICA) 29 May 1992	37,38, 40,43
Υ.	see page 14, line 26 - page 24, line 12; claims 1-20	1,3-8, 10-12, 26-30,35
,x	WO,A,9 305 178 (DAIKIN INDUSTRIES) 18 March 1993 cited in the application	37,38, 40,43,45
, ү	see page 29, line 1 - page 34, line 17; claims 1-59	1,3-8, 10-12, 26-30,35
i	· · · · · · · · · · · · · · · · · · ·	1
- 1	-/	1

INTERNATIONAL SEARCH REPORT

Into national application No.

PCT/US 93/03868

Box 1 Observations where certain claims were found unsearchable (Continueston of item 1 of first sheet)

This secure transfer of search report for not been established in respect of certain changi under Article 17(2)(a) for the following reasons: t. X Claum Nos. because they relate to subject matter not required to be searched by this Anthorny, namely: Remark: Although claims I-36 as far as they concern an "in vivo" treatment and 45,46 are directed to a method of treatment of the human/animal body the search has been carried out and based on the alleged effects of the compound/composition. Claims Nos.: because they relate to parts of the interm an extent that no meanineful internations nd are not drafted in accordance with the second and third seatences of Ruje 6.4(a). Bux II Observations where unity of invention is lacking (Continuation of item 2 of first sheet) This International Searching Authority fun As all required additional search fees were timely paid by the applicant, this international search report covers all old be searches without effort justifying an additional fee, this Authority did not invite payment As only some of the required additional search fees were timely paid by the applicant, this international search report covers only those claims for which fees were paid, specifically claims Noz.: No required additional search fees were timely paid by the applica-Remark on Protes The additional search feer were accompanied by the applicant's protest. No protest accompanied the payment of additional search fees.

ANNEX TO THE INTERNATIONAL SEARCH REPORT ON INTERNATIONAL PATENT APPLICATION NO.

US 9303868 SA 73747

This arrors lists the patent family members relating to the patent documents cited in the above-mentioned international search report.
The numbers are as continued in the European Patent Office EDP file or
Direspond Patent Office is in no way liable for them particulates which are merely given for the purpose of information.

09/07/93.

Patent document cited in search report	Publicacion date	Pace	Publication date	
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WO-A-9208791	29-05-92	AU-A-	9111691	11-06-92
WO-A-9305178	18-03-93	CA-A- EP-A- JP-T- WO-A-	2056983 0481065 4507198 9117267	08-11-91 22-04-92 17-12-92 14-11-91
WO-A-9220808	26-11-92	χη-γ-	2140292 0539573	30-12-92 05-05-93